

## INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 7:

A01N 63/00, 63/02, C12N 15/31, C12P 21/00, C07K 14/24 // (C12P 21/00, C12R 1:01)

(11) International Publication Number:

WO 00/42855

(43) International Publication Date:

27 July 2000 (27.07.00)

(21) International Application Number:

PCT/GB00/00219

(22) International Filing Date:

24 January 2000 (24.01.00)

(30) Priority Data:

9901499.5

22 January 1999 (22.01.99)

GB

(71) Applicant (for all designated States except US): HORTICUL-TURE RESEARCH INTERNATIONAL [GB/GB]; Wellesbourne, Warwick, Warwickshire CV35 9EF (GB).

(72) Inventors; and

(75) Inventors/Applicants (for US only): MORGAN, James, Alun, Wynne [GB/GB]; Horticulture Research International, Wellesbourne, Warwick, Warwickshire CV35 9EF (GB). JARRETT, Paul [GB/GB]; Horticulture Research International, Wellesbourne, Warwick, Warwickshire CV35 9EF (GB). ELLIS, Debbie [GB/GB]; Horticulture Research International, Wellesbourne, Warwick, Warwickshire CV35 9EF (GB). OUSLEY, Margaret, Anne [GB/GB]; Horticulture Research International, Wellesbourne, Warwick, Warwickshire CV35 9EF (GB).

(74) Agent: RUFFLES, Graham, Keith; Marks & Clerk, 57-60 Lincoln's Inn Fields, London WC2A 3LS (GB). (81) Designated States: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU, CZ, DE, DK, DM, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).

#### **Published**

With international search report.

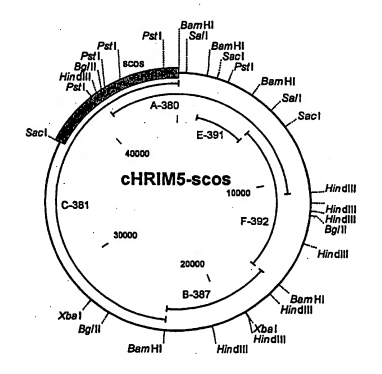
Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.

With an indication in relation to deposited biological material furnished under Rule 13bis separately from the description.

#### (54) Title: BIOLOGICAL CONTROL OF NEMATODES

#### (57) Abstract

Nematodes can be controlled through the use of bacteria associated symbiotically with an entomopathogenic nematode. The bacteria can be employed for nematode control, or engineered to a recombinant form. Control may be achieved using material such as a peptide. The peptide can be obtained from a natural or engineered nucleic acid.



# FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AL AM AT AU	Albania Armenia Austria	ES F1	Spain	LS	Lesotho		
AT		FI					
1	Austria		Finland	LT	Lithuania	SI	Slovenia
AII		FR	France	. LU	Luxembourg	SK	Slovakia
	Australia	GA	Gabon `	LV	Latvia	SN	Senegal
AZ	Azerbaijan	GB	United Kingdom	MC	Monaco	SZ	Swaziland
BA	Bosnia and Herzegovina	GE	Georgia	MD	Republic of Moldova	TD	Chad
BB	Barbados	GH	Ghana	MG	Madagascar	TG	Togo
BE	Belgium	GN	Guinea	MK	The former Yugoslav	TJ	Tajikistan
BF	Burkina Faso	GR	Greece	1,775	Republic of Macedonia	TM	Turkmenistan
BG	Bulgaria	HU	Hungary	ML	Mali	TR	Turkey
BJ	Benin	IE	Ireland	MN	Mongolia	TT	Trinidad and Tobago
BR	Brazil	IL	Israel	MR	Mauritania	UA	Ukraine
BY	Belarus	IS	Iceland	MW	Malawi	UG	Uganda
	Canada .	TI	Italy	MX	Mexico	US	United States of America
	Central African Republic	JP	Japan	NE	Niger	UZ	Uzbekistan
CG	Congo	KE	Kenya	NL	Netherlands	VN	Viet Nam
СН	Switzerland	KG	Kyrgyzstan	NO	Norway	YU	Yugoslavia
CI	Côte d'Ivoire	KP	Democratic People's	NZ	New Zealand	zw	Zimbabwe
СМ	Cameroon		Republic of Korea	PL	Poland		
CN	China	KR	Republic of Korea	PT			
	Cuba	ΚZ	Kazakstan	RO	Portugal Romania		
CZ	Czech Republic	LC	Saint Lucia	RU	Russian Federation		
DE	Germany	u	Liechtenstein	SD	Sudan		
	Denmark	LK	Sri Lanka	SE	Sweden		
EE I	Estonia	LR	Liberia	SG			
				30	Singapore		

### BIOLOGICAL CONTROL OF NEMATODES

TECHNICAL FIELD

The present invention relates to methods and materials for controlling nematodes.

PRIOR ART

Several thousand species of nematodes, sometimes called eel worms, are known. Numerous nematodes attack and parasitize humans and animals and cause disease. Additionally, several hundred species are known to feed on living plants. Certain of these are reviewed by Agrios in "Plant Pathology - 3rd Ed" Pub Academic Press Inc, see Chapter 15 therein.

Methods of controlling nematodes and their associated diseases include cultural practices; biological methods, e.g. use of resistant varieties; physical methods, e.g. heat; and use of chemical agents.

Patent application WO 92/19739 (Mycogen) relates to genes and gene fragments from *Bacillus thuringiensis* which have nematocidal activity. These generally encode crystal toxins from particular strains.

Patent application EP 0 303 426 (Mycogen) also relates to strains of *B. thuringiensis* which have nematocidal activity.

Patent application EP 0 171 381 (Monsanto) relates to particular soil bacteria which are capable of proliferating in an environment which is infested with

nematodes such as pseudomonads which colonise the surface of plant roots. The basis for the controlling activity appears to stem from glycosidase enzymes which are hypothesised to directly inhibit the nematodes.

Notwithstanding these disclosures, there is an ongoing requirement for materials which have nematocidal activity, for instance for use in crop protection or nematode-mediated disease control.

Patent application PCT/WO 99/22598 (University of Reading) published 14 May 1999 claims a biopesticide for the control of insect pests or plant parasitic nematodes or both, which comprises as an effective agent a species of bacteria which is a symbiont of an entomopathogenic nematode.

#### DISCLOSURE OF THE INVENTION

The present inventors have established that species of bacteria which in nature are associated symbiotically with entomopathogenic nematodes, can in fact be utilised to control nematodes, and in preferred forms of the invention, to kill them. The bacteria themselves can be employed, or nematode control agents can be used which are derived from such bacteria. In one aspect of the invention, the present invention employs bacteria which are engineered and thus not naturally occurring, or nematode control agents which are derived from natural or non-natural bacteria.

It has been reported that certain bacterial species such as *Xenorhabdus* and *Photorhabdus* can be used to control insects, see e.g. PCT/WO 98/08388 of MAFF, PCT/WO 97/17432 of WARF, and PCT/WO 99/42589 of Novartis. An effect against nematodes had not previously been demonstrated.

WO 00/42855 PCT/GB00/00219

3

The symbiotic bacteria used in the present invention are isolatable from nematodes or the insects which the nematodes attack, and differ fundamentally in terms of life-style and activity from those soil bacteria such as *B. thuringiensis* or pseudomonads which have previously been suggested as being nematocidal.

Indeed, prima facie, it seems highly unlikely that nematode symbiotes might possess nematocidal activity. However, in the light of the present disclosure, a number of possible explanations for the observed activity can be tentatively proposed. Firstly, in order to protect a nutrient supply from a dead insect, the bacteria might produce anti-nematocides to prevent saprophytic nematodes gaining access. Alternatively, to become a symbiont, the bacterial strains may have once been pathogens of these nematodes and evolved towards a less hostile symbiotic relationship. The nematocidal activity may be an evolutionary throwback from the original pathogenic relationship, in which case it may be expected to be widely present amongst bacteria which have evolved in this way.

A first aspect of the present invention is the use of bacterial strains to control a target nematode, characterised in that in nature the bacterial strain is associated symbiotically with an entomopathogenic nematode.

As discussed in more detail below, the bacterial strains may be used in the methods of the present invention *per se*, or they may be used as a source of nematode control agent. The nematode control agent can be derived directly, or be prepared and utilised through recombinant DNA techniques, optionally via a host cell.

The target nematode will generally be different to the nematode with which the bacterial strain is found symbiotically in nature.

By means of the present invention employing bacteria or a nematode control agent, it becomes possible to control nematodes, in the sense of, to prevent or retard the effect that the nematode has on other organisms such as animals or more preferably plants, or to reduce the number of nematodes or nematode eggs in an area of interest, or to alleviate or cure a disease caused by nematodes. Control may be at the level of larval nematodes or nematode eggs, or may inhibit the motion, feeding or infectivity of adult nematodes. Nematocidal control may be employed to kill the nematode target. Such controlling activity can be assessed as shown in the Examples below.

#### PREFERRED EMBODIMENTS

The present invention provides a composition for the control of parasitic nematodes which comprises as an effective agent a species of bacteria which is a symbiont of an entomopathogenic nematode, or engineered bacteria having such activity, or a nematode control agent derived from natural or engineered bacteria.

Correspondingly, the present invention also provides a method of nematode control employing such a composition.

The bacterial species is typically of the genera *Xenorhabdus* or *Photorhabdus*, preferably the genus *Xenorhabdus*, for instance the species *Xenorhabdus* bovienii. Examples of particularly preferred bacteria include:

Xenorhabdus bovienii strain H31 deposited with NCIMB under accession number NCIMB 40985 on 20 January 1999;

Xenorhabdus bovienii strain I73 deposited with NCIMB under accession number NCIMB 40986 on 05 November 1998; and

Xenorhabdus strain C42 deposited with NCIMB under accession number NCIMB 41004 on 05 November 1998.

The nematode control agent can be a peptide derived from a symbiont of an entomopathogenic nematode or an engineered bacterium has functional activity against a nematode. The peptide nematode control agent can be produced from a nucleic acid derived from a symbiont of an entomopath nematode or an engineered bacterium and which encodes such a peptide. The peptide can be an oligopeptide or a polypeptide, notably a protein. In one version, the nematode control agent is a toxin with toxic activity against nematodes, but the nematode control agent can have other activity.

The nucleic acids of this invention can be employed in a method of producing a peptide comprising the step of causing or allowing the expression from a nucleic acid of this invention in a suitable host cell.

The nucleic acid can comprise a natural nucleotide sequence or a degeneratively equivalent sequence, and functional variants thereof. Variants include homologous variants encoding a peptide which is a nematode control agent, the nucleic acid having 70% or more DNA sequence identity and/or the peptide having 70% or more amino acid sequence identity. Especially preferred nucleic acids in p 13-1f and p 14-2f and variants thereof.

The present invention extends to nucleic acids having a sequence which is a derivative by way of addition, insertion, deletion or substitution of one or more nucleotides. The nucleic acid can contain longer expressed sequences such that the nematode control agent is expressed as a fusion protein.

Nucleic acids complementary to the nucleic acid encoding a nematode

control agent are also part of this invention.

Nucleic acids for use as a probe or primer having a nucleotide sequence of at least 15, 18, 21, 24 or 30 nucleotides, which sequence is present in, or complementary to, the nucleic acid encoding nematode control agent are further provided by this invention. In this respect, the invention extends to a method for identifying or cloning a nucleic acid for nematode control agent which method employs such a nucleic acid probe.

A method provided by this invention comprises the steps of:

- (a) providing a preparation of nucleic acid from a bacterium,
- (b) providing a probe,
- (c) contacting nucleic acid in said preparation with said probe under conditions for hybridisation of probe to any said gene or homologue in said preparation, and,
- (d) identifying said gene or homologue if present by its hybridisation with said probe.

The hybridisation conditions can be selected to allow the identification of sequences having 70% or more sequence identity with the probe.

In one embodiment, the method comprises use of two primers to amplify a nucleic acid encoding a nematode control agent, at least one of the primers having a conserved nucleotide sequence of at least 15 nucleotides.

A method is further made possible by this invention comprising the steps of:

- (a) providing a preparation of nucleic acid from a bacterium,
- (b) providing a pair of nucleic acid molecule primers, at least one of which is a primer,
- (c) contacting nucleic acid in said preparation with said primers under

WO 00/42855 PCT/GB00/00219

7

conditions for performance of PCR,

(d) performing PCR and determining the presence of absence of an amplified PCR product.

Additionally, the invention provides a recombinant vector comprising a nucleic acid of this invention. The vector is preferably capable of replicating in a suitable host such as *E. coli* or in *Xenorhabdus*. The vector can be a baculovirus. In a preferred feature, the nucleic acid is operably linked to a promoter or other regulatory element for transcription in a host cell. Vectors can further comprise any one or more of the following: a terminator sequence; a polyadenylation sequence; an enhancer sequence; a marker gene; a sequence encoding pesticidal material derived from *Bacillus thuringiensis*.

The vector can be a plant vector.

The vector of this invention can be introduced into a cell. Thus, a method for transforming a plant cell comprises the step of causing or allowing recombination between the vector and the plant cell genome to introduce the nucleic acid into the genome. The nucleic acid can be incorporated into chloroplast DNA, or into mitochondrial DNA.

Host cells comprising a vector are also part of this invention. The host cell can be a plant cell, which may be in a plant.

To this end, a method for producing a transgenic plant comprises the step of regenerating a plant from the transformed cell. In turn, plants of this invention extend to the progeny of such plants.

Examples of plants of this invention include crop species which can be

protected, notably maize, cotton, soya, rice, *Brassica* species, tomato, potato, sugar beet, barley, soybean, peanut, onion, rye, wheat, corn, banana, raspberry, bean. Decorative and other plants are also possible, e.g. rose.

A part of the propagule of the plants is also envisaged by this invention.

A method of influencing or affecting the toxicity of a cell such as a plant cell is provided where the method includes causing or allowing expression of a heterologous nucleic acid of this invention within the cells.

In a further aspect, the invention involves the use of a material selected from: an *X. bovienii* strain, a nematode control agent; a nucleic acid; a host cell; a plant; a peptide; or a composition of the invention, for the control of a pest, especially where the pest is a nematode and the material is used to control the nematode.

The present invention extends to control of helminthiasis in humans and other animals including domesticated animals such as swine, sheep, horses, cattle, goats, dogs, cats and poultry. The nematodes to be controlled include Haemonchus, Trichostrongylus, Ostertagia, Nematodirus, Cooperia, Ascaris, Bunostomum, Oesophagosromuni, Chaberria, Trichuris, Strongylus, Trichonema, Dictyocaulus, Capillaria, Heterkis, Toxocara, Ascaridia, Oxyuris, Ancylostoma, Uncinaria, Toxascaris, Caenorhabditis and Parascaris.

Target nematodes may be selected from the genera Aphelenochoides,
Anguina, Bursaphalenchus, Criconemella, Melodigyne, Ditylenchus, Globodera,
Heliocotylenchus, Heterodera, Pratylenchus, Radopholus, Rotelynchus,
Tylenchus, Trichodorus, Xiphenema, and Caenorhabditis.

The compositions of this invention can be used in conjunction with Bacillus

thuringiensis or pesticidal materials derived therefrom.

In a further aspect, there is provided an antibody or fragment thereof, or a polypeptide comprising the antigen-binding domain of the antibody, capable of specifically binding a peptide of this invention.

Such an antibody or fragment can be obtained by immunising a mammal with the peptide, and is useful in a method of identifying and/or isolating a nematode control agent comprising the step of screening candidate polypeptides with a polypeptide comprising the antigen-binding domain of the antibody of claim.

Some further aspects of preferred embodiments of the invention will now be discussed.

## Bacterial strains

These can be derived from any entomopathogenic nematode. Preferred species are Xenorhabdus and Photorhabdus.

Potential sources of bacteria for use in the methods of the present invention may be identified by any preferred method. For instance, entomopathogenic nematodes can be isolated using an insect baiting technique such as that described by Bedding & Akhurst (1975) Nematologia 21: 215-227. Bacteria from nematodes identified as being pathogenic to the insect are isolated, cultured, and used as a source of nematocidal agent, e.g. by analogy with the methods used in the Examples below. Preferably Xenorhabdus or Photorhabdus species are used.

The preferred bacterial strains include ones which have the characteristics of

strain C42, I73 or H31 isolated by the present inventors. This *Xenorhabdus* strain has the following characteristics: rod shaped; motile; non-bio luminescent; blue on NBTA; produces antibiotics; resistant to ampicillin; forms circular colonies; has convex morphology; white colour.

This strain was presumptively identified as belonging to the genera *Xenorhabdus* since it was isolated from an insect killed by an entomopathogenic nematode and had the above characteristics. The strain has been deposited at the NCIMB (23 St Machar Drive, Aberdeen, AB24 3RY, Scotland) by the applicants under accession number NCIMB 41004 on 20 January 1999.

Further preferred strains of the present invention are two strains of X. bovienii designated H31 and I73 which have also been deposited under the terms of the Budapest Treaty at the NCIMB under the accession numbers NCIMB 40985 and 40986 respectively. These share characteristics of C42 in that they are rod-shaped; motile; non-bioluminescent; blue on NBTA; produce antibiotics; resistant to ampicillin; form circular colonies; and have convex morphology. The strains were identified as belonging to the species X. bovienii when compared to the X. bovienii type strain T228 using Restriction Analysis of the complete 16S rRNA gene and partial sequence analysis.

#### Target nematodes and diseases

The group of diseases described generally as helminthiasis is due to infection of an human or other animal host with parasitic worms known as helminths. Helminthiasis is a prevalent and serious economic problem in domesticated animals such as swine, sheep, horses, cattle, goats, dogs, cats and poultry. Among the helminths, the group of worms described as nematodes causes

WO 00/42855 PCT/GB00/00219

11

widespread and often at times serious infection in various species of animals. The most common genera of nematodes infecting the animals referred to above are Haemonchus, Trichostrongylus, Ostertagia, Nematodirus, Cooperia, Ascaris, Bunostomum, Oesophagosromuni, Chaberria, Trichuris, Strongylus, Trichonema, Dictyocaulus, Capillaria, Heterakis, Toxocara, Ascaridia, Oxyuris, Ancylostoma, Uncinaria, Toxascaris, Caenorhabditis and Parascaris. Certain of these, such as Nematodirus, Cooperia, and Oesophagostomum, attack primarily the intestinal tract, while others, such as Dictyocaulus are found in the lungs. Still other parasites may be located in other tissues and organs of the body.

The bacteria and encoded toxins of the invention may be used as nematocides for the control of the nematodes and diseases discussed above. More preferably, however, they are used to control soil and plant parasitic nematodes. Particular crop species which can be protected include tomatoes, potatoes, sugar beet, barley, soybean, peanut, onion, rye, wheat, corn, banana, raspberry, beans. Decorative and other plants may also be treated e.g. rose.

Target nematodes may be selected from the genera Aphelenochoides,
Anguina, Bursaphalenchus, Criconemella, Melodoigyne, Ditylenchus,
Globodera, Helicotylenchus, Heterodera, Pratylenchus, Radopholus,
Rotelynchus, Tylenchus, Trichodorus, Xiphenema. A further organism used
in certain of the Examples below is Caenorhabditis elegans. Other target
organisms and plants are discussed by Agrios in "Plant Pathology - 3rd Ed"
Pub Academic Press Inc, see Chapter 15 therein.

As stated above, the target nematode will generally be different to that with which the bacterial strain is found in nature.

Methods of use of bacteria

The bacteria may be used in any appropriate method which brings them into contact with the target nematode, preferably such that they, or their products, are ingested or absorbed by the target nematode.

In particular, regarding plants, the bacteria may be formulated in a variety of ways so as to enhance stability. For instance they may be employed in admixture with substrates to protect the cells.

The mixture can be spread over, ploughed into or otherwise mixed with nematode infected or potentially infected soil.

Regarding animals, bacteria intended for enteric inoculation can be mixed with carrier material that is suitable for ingestion by the intended animals.

Isolation of agent

Nematode control agents of the present invention, which may be proteinaceous, or nucleic acids encoding them, may be isolated and/or purified from the C42, I73 or H31 bacteria described above, in substantially pure or homogeneous form, or free or substantially free of other materials from the bacterial strain of origin. Where used herein, the term "isolated" encompasses all of these possibilities.

Methods of purifying proteins from heterogenous mixtures are well known in the art, e.g. selective precipitation, proteolysis, ultrafiltration with known molecular weight cut-off filters, ion-exchange chromatography, gel filtration, etc. A particularly useful initial technique in this regard is ultracentrifugation. Further methods which are known to be suitable for protein purification are disclosed in "Methods in Enzymology Vol 182 - Guide to Protein Purification" Ed. M P Deutscher, Pub. Academic Press Inc. Other references which outline techniques commonly used by those of ordinary skill in the art include "Protein Purification - principles and practice" Pub. Springer-Verlag, New York Inc (1982), and by Harris & Angal (1989) "Protein purification methods - a practical approach " Pub. O.U.P. UK.

Nematocidal activity may be assessed using a spread assay as discussed below.

The C42, I73 or H31 agent may be wholly or partially synthetic. In particular they may be recombinantly produced from nucleic acid sequences which are not found together in nature (do not run contiguously) but which have been ligated or otherwise combined artificially.

For instance, in the Examples below, nucleic acid encoding toxin(s) from I73 has been expressed in hosts cells using a vector system. Amino acid sequences of 38 different putative I73 toxin(s) are set out in sequence Annex I. These sequences are based on the nucleic acid sequence set out in Fig 2 ('chrim5'), a cosmid clone derived from I73 genomic DNA which conferred nematocidal activity upon *E. coli* cells into which it was introduced (i.e. significantly reduced nematode larval growth and development, and feeding). As detailed below, the entire amino acid sequence as set out in each case may not be required for nematocidal activity. In particular the portion up to the first Met in each sequence may be omitted, as may other portions which may not contribute to the nematocidal activity. Thus, not all the proteins or genes may be required for nematocidal activity, and usually there will be one or more principal proteins, though others may play supporting roles such as in enhancing the activity or encoding other nematocidal activities.

Thus isolated nematocidal agents comprising a polypeptide containing all, or a nematocidal fragment, of any of the depicted I73 sequences, form one aspect of the present invention. Preferred agents include those encoded by p14-2f and p13-1f. Other active variants of these sequences are also encompassed as described below.

Candidate agents for use in this invention to control nematodes extend to those from the bacteria described in PCT/WO 99/22598, as well as the insecticidal toxins and bacteria of PCT/WO 99/42589, PCT/WO 98/08388 and PCT/WO 97/17432, the disclosures of which are incorporated by reference.

#### Nucleic acids and variants

In one aspect of the present invention there is provided a nucleic acid molecule encoding a nematode control agent of the present invention, for example a toxin, as described above.

The nucleic acid may be derived from the sequence shown in Fig 2 or the complement (or degenerate equivalent) thereof. This sequence (cHRIM5) was itself derived from I73 and identified by its unexpected nematocidal activity. Regions of this sequence believed to correspond to genes of the present invention are described in Fig 3. Isolated nucleic acids comprising one or more of these regions which encode a nematocidal activity are particularly preferred.

In the light of the present disclosure, further nucleic acids of the present invention may be isolated using PCR or southern blotting or other techniques well known to those skilled in the art. This requires the use of two primers to specifically amplify target nucleic acid, so preferably two

nucleic acid molecules with sequences characteristic of the C42, H31 or most preferably an I73 toxin isolated as above are employed. Using RACE PCR, only one such primer may be needed (see "PCR protocols; A Guide to Methods and Applications", Eds. Innis et al, Academic Press, New York, (1990)).

Thus a method involving use of PCR in obtaining nucleic acid according to the present invention may include:

- (a) providing a preparation of bacterial nucleic acid,
- (b) providing a pair of nucleic acid molecule primers suitable for PCR, at least one of said primers being a primer based on a toxin from C42, H31 or I73,
- (c) contacting nucleic acid in said preparation with said primers under conditions for performance of PCR,
- (d) performing PCR and determining the presence or absence of an amplified PCR product. The presence of an amplified PCR product may indicate identification of a variant.

In a further aspect of the present invention there are disclosed nucleic acids which are variants of the C42, I73 or H31 toxin. A variant nucleic acid molecule shares homology (or identity) with all or part of the C42, H31, or most preferably I73 sequence discussed above.

Preferably sequence comparisons are made using FASTA and FASTP (see Pearson & Lipman, 1988. Methods in Enzymology 183: 63-98). Parameters are set, using the default matrix blosum62, as follows:

Gapopen (penalty for the first residue in a gap): -12 for proteins / -16 for DNA

Gapext (penalty for additional residues in a gap): -2 for proteins / -4 for DNA KTUP word length: 2 for proteins / 6 for DNA.

Homology (similarity or identity) may be at the nucleotide sequence and/or encoded amino acid sequence level. Preferably, the nucleic acid and/or amino acid sequence shares at least about 70%, 75%, 80%, or 85% homology, most preferably at least about 90%, 95%, 96%, 97%, 98% or 99% homology.

Another method for assessing homology at the nucleic acid level is by hybridization screening. One common formula for calculating the stringency conditions required to achieve hybridisation between nucleic acid molecules of a specified sequence homology is shown in Molecular Cloning: a Laboratory Manual: 2nd edition, Sambrook et al, 1989, Cold Spring Harbor Laboratory Press:

Tm = 81.5°C + 16.6Log [Na+] + 0.41 (% G+C) - 0.63 (% formamide) - 600/#bp in duplex

As an illustration of the above formula, using [Na+] = [0.368] and 50-% formamide, with GC content of 42% and an average probe size of 200 bases, the Tm is 57°C. The Tm of a DNA duplex decreases by 1 - 1.5°C with every 1% decrease in homology. Thus, targets with greater than about 75% sequence identity would be observed using a hybridization temperature of 42°C. Such a sequence would be considered substantially homologous to the nucleic acid sequence of the present invention.

Variants of the present invention can be artificial nucleic acids.

Alternatively they may be novel, naturally occurring, nucleic acids, isolatable using the information disclosed herein. Thus a variant may be a distinctive part or fragment (however produced) corresponding to a portion of the C42, I73 or H31 toxin. The fragments may encode particular functional parts of the agent or they may be used for probing for, or amplifying, sequences corresponding to C42, I73 or H31 toxin. Sequence variants which occur naturally may include homologs of the C42, I73 or H31 toxin from other

In one aspect of the present invention, the nucleic acid encoding the nematode control agent is provided in the form of a recombinant and preferably replicable vector.

Generally speaking, those skilled in the art are well able to construct vectors and design protocols for recombinant gene expression. Suitable vectors can be chosen or constructed, containing appropriate regulatory sequences, including promoter sequences, terminator fragments, polyadenylation sequences, enhancer sequences, marker genes and other sequences as appropriate. For further details see, for example, Sambrook et al (1989) supra.

The permitted vectors include, *inter alia*, any plasmid, cosmid, phage or Agrobacterium binary vector in double or single stranded linear or circular form which may or may not be self transmissible or mobilizable, and which can transform a prokaryotic or eukaryotic host either by integration into the cellular genome or exist extrachromosomally, e.g. an autonomous replicating plasmid with an origin of replication. Illustratively integration can occur into chloroplast DNA or into mitochondrial DNA.

Preferably the nucleic acid in the vector is under the control of, and operably linked to, an appropriate optionally inducible promoter or other regulatory elements for transcription in a host cell such as a microbial, e.g. bacterial, yeast, filamentous fungal or plant cell. The vector may be a bi-functional expression vector which functions in multiple hosts. In the case of genomic DNA, this may contain its own promoter or other regulatory elements and in the case of cDNA this may be under the control of an appropriate promoter or other regulatory elements for expression in the host cell. The vectors and host cells into which they are introduced may be used to clone or otherwise

identify nucleic acids according to the invention.

The agent may be used as part of a viral vector which is itself pathogenic to nematodes.

Also of interest in the present context are nucleic acid constructs which operate as plant vectors. Specific procedures and vectors previously used with wide success upon plants are described by Guerineau and Mullineaux (1993) (Plant transformation and expression vectors. In: Plant Molecular Biology Labfax (Croy RRD ed) Oxford, BIOS Scientific Publishers, pp 121-148). Suitable vectors may include plant viral-derived vectors (see e.g. EP-A-194809). Suitable promoters which operate in plants include the Cauliflower Mosaic Virus 35S (CaMV 35S). Other examples are disclosed at page 120 of Lindsey & Jones (1989) "Plant Biotechnology in Agriculture" Pub. OU Press, Milton Keynes, UK.

## Host cells

The toxin genes or gene fragments encoding the nematocidal agents of the subject invention may be introduced into a host cell, microbial, animal or plant. Expression of the toxin gene in the host cell results, directly or indirectly, in the intracellular production and maintenance of the nematocide.

Thus the present invention also provides methods comprising introduction of such a construct into a plant cell or a microbial cell and/or induction of expression of a construct within a cell, by application of a suitable stimulus e.g. an effective exogenous inducer.

Hosts may be used to assay the activity of particular sequences or

fragments. Hosts can also be used to generate quantities of toxin which can be employed in situ in suitable treated cells, or alternatively with suitable hosts, e.g., *Pseudomonas* viable microbes can be applied to the sites of nematodes where they will proliferate and where they or their products can be ingested by the nematodes. Higher organisms, preferably plants, can also be engineered with the toxin. The result in each case is a control of the nematodes. A host may be selected that can tolerate harsh environmental conditions and then grow when they improve, as illustrated by *Bacillus* species where the spores can exist under environmental extremes.

Characteristics of interest for use as a nematocide microcapsule i.e. a vehicle for the active agent include protective qualities for the nematocide, such as thick cell walls, pigmentation, and intracellular packaging or formation of inclusion bodies; leaf affinity; lack of mammalian toxicity; attractiveness to nematodes for ingestion; ease of killing and fixing without damage to the toxin; and the like.

#### Treated host cells

Where the cell is treated, the cell will usually be intact and be substantially proliferative form when treated, rather than in a spore form, although in some instances spores may be employed. Treatment of the microbial cell, e.g. a microbe containing the bacterial toxin gene or gene fragment, can be by chemical or physical means, or by a combination of chemical and/or physical means, so long as the technique does not deleteriously affect the properties of the toxin, nor diminish the cellular capability in protecting the toxin.

## Viable hosts

Where the toxin gene or gene fragment is introduced via a suitable vector into a microbial host, and said host is applied to the environment in a living state, it is preferable that microorganism hosts are selected which are known to occupy the phytosphere (phylloplane, phyllosphere, rhizosphere, and/or rhizoplane) of one or more crops of interest. These microorganisms are selected so as to be capable of successfully competing in the particular environment (crop and other insect habitats) with the wild-type microorganisms, provide for stable maintenance and expression of the gene expressing the polypeptide pesticide, and, desirably, provide for improved protection of the nematocide from environmental degradation and inactivation.

A large number of microorganisms are known to inhabit the phylloplane (the surface of the plant leaves) and/or the rhizosphere (the soil surrounding plant roots) of a wide variety of important crops. These microorganisms include bacteria, algae, and fungi. Of particular interest are microorganisms, such as bacteria, e.g., genera Pseudomonas, Erwinia, Serratia, Klebsiella, Xanthomonas, Streptomyces, Rhizobium, Rhodopseudomonas, Methylophilius, Agrobacterium, Acetobacter, Lactobacillus, Arthrobacter, Azotobacter, Leuconosroc, and Alcaligenes; fungi, particularly yeast, e.g., genera Saccharomyces, Cryprococcus, Kluyveromyces, Sporobolomyces, Rhodororula, and Aureobasidium.

#### Plants as hosts

Nucleic acid encoding the nematocides of the present invention can be introduced into plant cells using any suitable technology, such as a disarmed Ti-plasmid vector carried by *Agrobacterium* exploiting its natural gene transfer ability (EP-A-270355, EP-A-0116718, NAR 12(22) 8711 - 87215 1984), particle or microprojectile bombardment (US 5100792, EP-A-444882.

EP-A-434616) microinjection (WO 92/09696, WO 94/00583, EP 331083, EP 175966, Green et al. (1987) Plant Tissue and Cell Culture, Academic Press), electroporation (EP 290395, WO 8706614 Gelvin Debeyser) other forms of direct DNA uptake (DE 4005152, WO 9012096, US 4684611), liposome mediated DNA uptake (e.g. Freeman et al. Plant Cell Physiol. 29: 1353 (1984)), or the vortexing method (e.g. Kindle, PNAS U.S.A. 87: 1228 (1990d). Physical methods for the transformation of plant cells are reviewed in Oard, 1991, Biotech. Adv. 9: 1-11.

Agrobacterium transformation is widely used by those skilled in the art to transform dicotyledonous species. It has also been used with filamentous fungi (see de Groot et al, 1998, Nature Biotechnology 16: 839-842).

Recently, there has also been substantial progress towards the routine production of stable, fertile transgenic plants in almost all economically relevant monocot plants (see e.g. Hiei et al. (1994) The Plant Journal 6, 271-282)). Microprojectile bombardment, electroporation and direct DNA uptake are preferred where *Agrobacterium* alone is inefficient or ineffective. Alternatively, a combination of different techniques may be employed to enhance the efficiency of the transformation process, e.g. bombardment with *Agrobacterium* coated microparticles (EP-A-486234) or microprojectile bombardment to induce wounding followed by co-cultivation with *Agrobacterium* (EP-A-486233).

Generally speaking, following transformation, a plant may be regenerated, e.g. from single cells, callus tissue or leaf discs, as is standard in the art. Almost any plant can be entirely regenerated from cells, tissues and organs of the plant. Available techniques are reviewed in Vasil et al., Cell Culture and Somatic Cell Genetics of Plants, Vol I, II and III, Laboratory Procedures and Their Applications, Academic Press, 1984, and Weissbach and

Weissbach, Methods for Plant Molecular Biology, Academic Press, 1989.

The generation of fertile transgenic plants has been achieved in the cereals rice, maize, wheat, oat, and barley (reviewed in Shimamoto, K. (1994) Current Opinion in Biotechnology 5, 158-162.; Vasil, et al. (1992) Bio/Technology 10, 667-674; Vain et al., 1995, Biotechnology Advances 13 (4): 653-671; Vasil, 1996, Nature Biotechnology 14 page 702).

#### Combination nematocides

In further embodiments of the invention, bacteria associated with entomopathogenic nematodes or the toxins or products discussed above are used in conjunction with other nematocidal bacteria such as *B. thuringiensis* strains (e.g. from WO 92/19739) or pesticidal materials derived therefrom.

Materials for use in the present invention

The present invention also embraces materials for use in the methods above. These materials include the novel bacterial strains which are associated symbiotically with an entomopathogenic nematode and which are capable of controlling a target nematode. In particular the invention encompasses strain C42, I73 or H31 in isolated or substantially isolated form, or strains having the characteristics of C42, I73 or H31 (including nematocidal activity assessed as below).

Also embraced are compositions and formulations of these bacteria. These may comprise or consist of wettable powders, granules or dusts, mixed with various inert materials, such as inorganic minerals (phyllosilicates, carbonates, sulfates, phosphates, methylcellulose, xanthan gum and the like) or botanical materials (powdered corncobs, rice hulls, walnut shells,

WO 00/42855 PCT/GB00/00219

24

peat moss, vermiculite, soil, seeds, other plant tissue and the like). The formulations may include spreader-sticker adjutants, stabilizing agents or surfactants. Liquid formulations may be aqueous-based or non-aqueous and employed as foams, gels, suspensions, emulsifiable concentrates, or the like. The ingredients may include rheological agents, surfactants, emulsifiers, dispersants, or polymers.

Bacteria may be mixed with other material while in freeze-dried form, encapsulated in biodegradable or water-soluble material, or otherwise treated to prolong their viability or decrease their levels of metabolic activity during handling. If desired, the carrier material may contain assimilatable nutrient sources to support proliferation of the bacteria.

Also included are purified or substantially purified nematocidal agents (particularly proteinaceous agents) isolated or isolatable from the strains or host cells discussed above.

Thus the invention further discloses nematocidal compositions comprising one or more agents as described above. Such compositions preferably further comprise other nematocidal materials from other *Xenorhabdus* species or non-*Xenorhabdus* species. These other materials may be chosen such as to have complementary properties to the agents described above, or act synergistically with it.

Toxins of the invention for use with animals can be adapted to be administered orally in a unit dosage form such as a capsule, bolus or tablet, or as a liquid drench when used as an anthelmintic in mammals, and in the soil to control plant nematodes. The drench is normally a solution, suspension or dispersion of the active ingredient, usually in water, together with a suspending agent such as bentonite and a wetting agent or like

excipient. Generally, the drenches also contain an antifoaming agent. Drench formulations generally contain from about 0.001 to 0.5% by weight of the active compound. Preferred drench formulations may contain from 0.01 to 0.1% by weight, the capsules and boluses comprise the active ingredient admixed with a carrier vehicle such as starch, talc, magnesium stearate, or dicalcium phosphate. Where it is desired to administer the toxin compounds in a dry, solid unit dosage form, capsules, boluses or tablets containing the desired amount of active compound usually are employed. These dosage forms are prepared by intimately and uniformly mixing the active ingredient with suitable finely divided diluents, fillers, disintegrating agents and/or binders such as starch, lactose, talc, magnesium stearate, vegetable gums and the like. Such unit dosage formulations may be varied widely with respect to their total weight and content of the antiparasitic agent, depending upon the factors such as the type of host animal to be treated, the severity and type of infection and the weight of the host.

When the active compound is to be administered via an animal feedstuff, it is intimately dispersed in the feed or used as a top dressing or in the form of pellets which may then be added to the finished feed or, optionally, fed separately. Preferably, a carrier for feed administration is one that is, or may be, an ingredient of the animal ration. Suitable compositions include feed premixes or supplements in which the active ingredient is present in relatively large amounts and which are suitable for direct feeding to the animal or for addition to the feed either directly or after an intermediate dilution or blending step. Typical carriers or diluents suitable for such compositions include, for example, distillers' dried grains, corn meal, citrus meal, fermentation residues, ground oyster shells, wheat shorts, molasses solubles, corn cob meal, edible bean mill feed, soya grits, crushed limestone and the like.

Alternatively, the antiparasitic compounds may be administered to animals parenterally, for example, by intraluminal, intramuscular, intratracheal, or subcutaneous injection, in which event the active ingredient is dissolved or dispersed in a liquid carrier vehicle. For parenteral administration, the active material is suitably admixed with an acceptable vehicle, preferably of the vegetable oil variety, such as peanut oil, cotton seed oil and the like. Other parenteral vehicles, such as organic preparations using solketal, glycerol, formal and aqueous parenteral formulations, are also used. The active compound or compounds are dissolved or suspended in the parenteral formulation for administration; such formulations generally contain from 0.005 to 5% by weight of the active compound.

Further aspects of the invention include nucleic acids, vectors and host cells containing a heterologous construct according to the present invention, especially a plant or a microbial cell.

Such microbial cells may be treated as described in the methods above. Examples of chemical reagents are halogenating agents. Other suitable techniques include treatment with aldehydes, such as formaldehyde and glutaraldehyde; anti-invectives, such as zephiran chloride and cetylpyridinium chloride; alcohols, such as isopropyl and ethanol; various histologic fixatives, such as Bouin's fixative and Helly's fixative (See: Humason, Gretchen L., Animal Tissue Techniques, W.H. Freeman and Company, 1967); or a combination of physical (heat) and chemical agents that preserve and prolong the activity of the toxin produced in the cell when the cell is administered to the host animal. The method of inactivation or killing retains at least a substantial portion of the bio-availability or bioactivity of the nematode control agent.

In all of the compositions discussed above, the nematocide concentration may vary widely depending upon the nature of the particular formulation, particularly whether it is a concentrate or to be used directly. The nematocide will be present in at least 1% by weight and may be 100% by weight. The dry formulations will have from about 1-95% by weight of the nematocide while the liquid formulations will generally be from about 16% by weight of the solids in the liquid phase. The formulations will generally have from about 10² to about 10¹0 cells/mg, more preferably 10² to about 10⁰ cells/mg. These formulations will be administered at about 50 mg (liquid or dry) to 1 kg or more per hectare. The formulations can be applied to the environment of the nematodes, e.g., plants, soil or water, by spraying, dusting, sprinkling, or the like.

In addition to the above the invention includes plant cells which have been transformed with the genes of the present invention, and plants which include such plant cells.

#### EXAMPLES OF THE INVENTION

The invention will now be further described with reference to the following non-limiting Figures and Examples. Other embodiments of the invention will occur to those skilled in the art in the light of these.

#### **FIGURES**

Fig 1 shows the cHRIM5 cosmid vector and subclones used for sequencing, as described in Example 6.

WO 00/42855 PCT/GB00/00219

28

Fig 2 shows the sequence of cHRIM5 (1-37544 bps).

Fig 3 shows the position and orientation of ORFs in the cHRIM5 sequence.

Fig 4 shows deletions of cHRIM5 tested for nematocidal activity.

Fig 5 illustrates cloning of nematocidal activity in PLEX.

## Example 1 - Source of strains C42, I73 and H31

Strain C42 was obtained using an insect entrapment method. Insects which were killed on the surface of a soil sample were observed under a microscope at high magnification. Any that contained high numbers of bacteria and not fungal hyphae were presumed to have been killed by insect parasitic nematodes. The identified presence of nematodes also aids this identification step, but it is not essential. These samples were plated on to NBTA media (see Poinar & Thomas, 1984 Nematodes p238-280 in "Laboratory guide to insect pathogens and parasites" Eds. Poiner & Thomas, Pub. Plenum Press, New York). Any colonies that developed that had characteristic features (e.g. morphology, size, colour) of *Xenorhabdus* or *Photorhabdus* strains were selected. Non-luminescent colonies were presumptively identified as *Xenorhabdus*. The identity of those having nematocidal activity as assessed in Example 3, is further confirmed using 16s rRNA sequence data (see Brunel et al 1997, Applied and Environmental Microbiology 63,2: 574-580).

I73 and H31 strains were obtained in a similar way to strain C42 but they were identified as belonging to the species *X. bovienii* when compared to the *X. bovienii* type strain T228 using Restriction Analysis of the complete 16S

rRNA gene (see Brunel et al, 1997 Applied and Environmental Microbiology: 574-580), and partial 16s ribosomal RNA sequence analysis.

## Example 2 - Cell growth and preservation

Subcultures of the *Xenorhabdus* species C42, I73 and H31 were used to inoculate three 9 cm diameter petri dishes containing L agar (10g tryptone, 5 g Yeast Extract, 5 g NaCl and 15 g agar per lt). Plates were incubated for 48 hrs at 26°C and the resulting growth harvested by scraping off bacterial cells and thoroughly resuspending in 40 mls of 5% w/v lactose. The cells were washed once by centrifugation (5000 x g for 10 mins), resuspended in 10 mls of 5% w/v lactose, dispensed into 1 ml aliquots and freeze dried (-60°C for 48 hrs ) for medium term storage at 2°C. Other stocks were resuspended in nutrient broth containing 10% w/v glycerol (Protect) and frozen at -70°C.

## Example 3- Activity of cells against Caenorhabditis elegans

The bioassays were performed by allowing *C. elegans* to feed on live bacterial cell suspensions spread over the surface of Luria broth agar (Luria broth containing 1.2%w/v agar) in segmented square petri dishes (2.0 x 2.0 cm per test well). A minimum of three test wells, each containing 50-100 nematodes were used for each test. Mortalities were recorded after 3 days at 18°C.

C. elegans was cultured on Escherichia coli at 18°C on 9 cm diameter LB agar plates. Once the nematodes had colonised the complete plate they were resubbed on a fresh plate to maintain stocks and the remainder re-suspended in 40 ml LB. The tube was allowed to stand for 15 min and the nematodes settled to the bottom. The concentrated nematodes were removed using a

PCT/GB00/00219

WO 00/42855

sterile pipette and placed in 40 mls of fresh LB. The process was repeated 5 more times to wash the nematodes away from the E. coli cells. The nematodes were then diluted so that approximately 50 nematodes were present in 50 µl of LB.

The Xenorhabdus cells used were cultured in LB at 30°C/100 rpm for 24 hours and 50  $\mu$ l spread on to the surface of each test well. The control E. coli cells were treated in a similar way but incubated at 37°C for growth. After application the wells were air dried for 30 min, and 50 µl of the nematode suspension placed in each well. Again the wells were air dried for 30 min. Plates were incubated at 18°C with 80% relative humidity for 3 days.

Xenorhabdus spp. C42, H31 and I73 gave 95% mortality, as compared with no significant effect for certain other Xenorhabdus bacterial strains and E. coli. Thus these results clearly show that cells from Xenorhabdus C42, H31 and I73 are an effective nematocide.

## Example 4 - Cloning of nematode active gene from I73

Total DNA was isolated from I73 using a Quiagen genomic DNA purification kit (cat no. 10243). To isolate DNA, cells were grown in Luria broth (10g tryptone, 5g yeast and 5g NaCl per lt) at 26°C with shaking at 200 rpm to an optical density of 1.5 A600. Cells were harvested by centrifugation at 4000 x g and the DNA isolated using Quiagen 100/G tips, as per manufacturer's instructions. The purified DNA was stored at -20°C in TE buffer (10 mM Tris, 1 mM EDTA, pH 8.0).

To obtain a representative 173 library, total DNA was partially digested with Sau3A. Approximately 25 µg of DNA was incubated at 37°C with 0.25 units of enzyme. At intervals of 5, 15, 30, 45 and 60 minutes, samples were removed and heated at 65°C for 10 minutes. To determine the size of the resulting DNA fragments, the samples were separated on a 0.5% (w/v) agarose gel. The samples containing a dominant DNA fragment size of between 30 and 50 Kb were combined and treated with shrimp alkaline phosphatase (Boehringer) for 20 minutes at 37°C. The DNA was ligated into the *Bam*HI site of the Stratagene cosmid vector Supercos1 (scos) and packaged into the *Escherichia coli* strain XL Blue 1, using a Gigapack II packaging kit (Stratagene) following the manufacturer's instructions.

To identify individual cosmid clones with activity to C. elegans, single colonies were grown in individual wells of segmented square petri dishes on Luria agar, containing 50  $\mu$ g/ml ampicillin at 30°C for 24 hours. To each well, approximately 50, mainly L4 and adult C. elegans larvae were added in 50  $\mu$ l of Luria broth. The dishes were incubated at 18°C and examined after 6 days for nematode development.

A total of 600 clones were examined and one coded cHRIM5 was found, which caused significant reduction in larval numbers, with no live L4 and adult larvae observed compared to on average, greater than 40 in all other clones tested.

## Example 5 - Activity of cHRIM5, C42, H31 and I73 against C. elegans

Clone cHRIM5 was grown in 50 mls LB containing 50  $\mu$ g of ampicillin per ml at 30°C/200 rpm for 40 hours. C42, H31 and I73 were grown in 50 mls LB at 26°C for 48 hours/200 rpm. Cultures were centrifuged at 4000 x g for 10 minutes, washed once and resuspended in 5 mls of PBS (0.05 mM phosphate buffer, 0.125M NaCl). To determine activity, 300  $\mu$ l of cells were added in triplicate, to 1.2 ml of PBS containing 25, mainly L4 and adult C.

elegans larvae in multi well square dishes. As a control, an equivalent amount of XL 1 Blue *E. coli* cells containing Supercos 1 were used to determine nematode survival. The assays were incubated at 18 °C for 7 days before approximate nematode counts and observations were made.

## Activity of cells on C. elegans

Cell line	No. and size of	Cell turbidity
	larvae/square	
XL 1 Blue/Supercos 1	>100 (all stages)	Clear
XL 1 Blue/cHRIM5	<20(mainly small, L1,2 &3)	Cloudy
C42	<10	Cloudy
H31	<10	Cloudy
173	<10	Cloudy

Thus cHRIM5, C42, H31 and I73 all gave a reduction in nematode numbers, and in particular cHRIM5 cells significantly reduced larval growth and development. All four strains caused a reduction in feeding (as indicated by the cloudy cell suspensions).

## Example 6 - DNA and protein sequences

Plasmid and cosmid DNA for cloning was prepared using the QIAGEN midi system (tip 100, cat. No 12143). Cells were grown in Luria broth (Merck) at  $37^{\circ}$ C with shaking at 200 rpm for 18 hours. Cells were harvested by centrifugation at  $6,000 \times g$  and the DNA isolated as per manufacturers instructions. Restriction digestion (Roche, Life Technologies), dephosphorylation (Roche) and ligation (Life Technologies) were carried out using manufacturer's recommended conditions and as outlined by Sambrook et al. Transformation was accomplished using electrocompetant cells and a

BIO-RAD Gene pulser set at 12.5V cm<sup>-2</sup>. Two  $\mu$ l of DNA was used to electroporate 80  $\mu$ l of early log phase *E. coli* DH5 alpha cells washed 3 times in sterile water (centrifugation at 6000 x g for 5 mins) and resuspended in 1/100th the original volume in 10% (v/v) glycerol. Luria agar containing either kanamycin or ampicillin at 50  $\mu$ g ml<sup>-1</sup> were used to select clones where appropriate.

DNA sequence analysis of cHRIM5 was completed by sequencing a number of sub clones and primer walking, see figure 1 for the supercos vector, where the numbers are kBp. The sub clones used are as follows:

code	cHRIM5 treatment	vector used or remaining
A-380	HindIII digestion and self-ligation	deleted scos
B-387	BamHI digestion and self-ligation	pUC 19-BamH1 digestion
C-381	Sall-BamHI digestion	scos
E-391	Sall-BamHI digestion	pUC 19-Sall BamH1 digestion
F-392	Sall-BamHI digestion	pUC 19-Sall BamH1 digestion

Sub clone A-380 was constructed by digesting cHRIM5 DNA with the restriction enzyme *Hind*III and re-ligating fragments, this clone contains a deletion of the insert and scos cosmid DNA as the vector. Sub clone B-387 is a *Bam*HI digestion of cHRIM5 cloned into the plasmid pUC19 also cut with *Bam*HI and dephosphorylated. Sub clone C-381 was obtained by digesting CHRIM5 DNA with *Bam*HI and re-ligating the fragments, this clone contains the scos cosmid as the vector. Clones E-391 and F-392 were obtained by cutting cHRIM5 DNA with *Sal*I and *Bam*HI and ligating these fragments into the vector pUC19 also cut with these enzymes.

Sequencing was conducted using the artificial transposon AT2 (supplied by Perkin-Elmer-Applied Biosystems, Primer Island Transposition kit, cat No.

403015) using the cosmid cHRIM5 and all sub-clones as target DNA. One μg of cHRIM5 DNA was incubated with the transposon AT2 for 1 hour at 30°C in a final volume of 20 µl. After incubation the reaction was stopped by adding 5 µl of 0.25M EDTA, 1% (w/v) SDS, and heat treatment at 65°C for 30 mins. The DNA was desalted by dialysis against water. One µl of the reaction mix was used to electroporate 80 µl of early log phase E. coli DH5 alpha cells. Colonies were selected on LB media containing 50 µg/ml trimethoprim. Once inserted the transposon mutants were used to provide a range of positions of primer sites at random intervals throughout the clones. The two primers PI+ and PI- near the end of the transposon were used to generate sequence data. In addition standard primers for the pUC19 and scos vectors were used to generate sequence data at the ends of each clone. DNA for sequencing was prepared using the QIAGEN ion exchange media (qiawell8, cat. No. 17122). Clones were grown in 1 ml of Luria broth containing trimethoprin (50 µg ml<sup>-1</sup>) for 18 hours. Cells were centrifuged at 13,000 x q for 5 mins and resuspended in 350 µl of buffer P1. After 5 mins 350 µl of buffer P2 was added and the samples incubated for 5 mins at room temperature. To this 350 µl of buffer P3 was added and the samples left on ice for 15 mins. After centrifugation at 13,000 x q for 15 mins the samples were loaded on the Qiagen column under vacuum, and washed with buffer QC. DNA was eluted with buffer QF (500 μl) at 50°C and isopropanol precipitated (0.8 vol). After centrifugation at  $13,000 \times g$  for 30min, DNA was washed with 70% (v/v) ethanol and air dried for 10 mins. The final pellet was resuspended in 10 µl of water. Cycle sequencing reactions using the Perkin-Elmer Applied Biosystems division Big Dye reaction kit (cat No. 4303149) were prepared using standard conditions for plasmid and cosmid sequencing. Samples were analysed on ABI Automated Sequencers. DNA sequences were assembled using the DNA\* software. The complete sequence of cHRIM5 was obtained by primer walking to join the final DNA contigs together. The final sequence of cHRIM5ed2 is shown

in Figure 2. Analysis of the DNA using the software Clone indicated a number of ORF illustrated in Figure 3 and 4. Corresponding protein sequences are also presented at Annex I.

## Example 7 - Fragments that encode nematocidal activity

To identify smaller fragments that encoded nematocidal activity, a series of sub-cloning experiments were performed using *E. coli* DH5 alpha. Qiagen midi and miniprep methods, restriction and ligations were used as for previous examples. Nematicidal activity of all constructs was determined as described in Example 4. In Figure 4, we show the deletions of cHRIM5 tested for nematicidal activity. Restriction sites and genes are indicated. Size in base pairs indicated on the map line. A cHRIM5, B cHRIM6, C cHRIM7, D cHRIM8, E cHRIM8, F cHRIM10, G *Ndel* deletion of cHRIM8, H Approximate positions (arrows) of three AT2 transposon insertions (tn58, tn26, tn43) in cHRIM9.

The cosmid cHRIM5 (figure 4A) was digested with the enzyme Sall and religated. The resulting sub clone cHRIM6, illustrated in Figure 4B showed nematicidal activity. cHRIM6 was digested with the enzyme Smal and religated, producing sub-clone cHRIM7 (Figure 4C). CHRIM7 was digested with BglII and the kanamycin resistance gene block (nptII, Pharmacia) cut with BamHI was ligated into it. After selection on LB containing kanamycin (50µg ml-1) and ampicillin (50µg ml-1) the clone was digested with Sall and religated, in effect creating a deletion from the Sall site to the BglII site of cHRIM6 to generate CHRIM8 (figure 4D). By cutting cHRIM8 with Nrul a further deletion was made to create cHRIM10 (figure 4F). All the above clones maintained nematicidal activity.

Deletion of cHRIM8 with Ndel, removed a portion of the p14-2f gene (figure

WO 00/42855 PCT/GB00/00219

36

4G), this reduced nematicidal activity. This indicates that the p14-2f gene or protein are important for nematocidal activity. Transposon mutagenesis of cHRIM9 (a clone very similar to cHRIM7 but deleted with Narl rather than Smal) with the artificial transposon AT2 (Perkin Elmer Applied Biosystems) resulted in a number of inserts within this clone (figure 4H). Insert cHRIM9-tn43 was restriction mapped to an approximate position of bp 20,700 (on cHRIM5) within the p20-9r gene, this mutant retained nematicidal activity. This indicates that this gene is not essential for activity. Insert cHRIM9-tn58 mapped to an approximate position of bp 13,400 (on cHRIM5), within the p13-1f gene, nematicidal activity was reduced. This indicates that this gene, region of DNA or the blocking effect of the transposon in this position is important for activity. Insert cHRIM9tn26 was restriction mapped to approximate position of bp 15,000 (on cHRIM5) within the p14-2f gene, nematocidal activity was reduced. This indicates that this gene, region of DNA or the blocking effect of the transposon in this position is important for activity.

Clone cHRIM6-tn43 was digested with *BgI*II and *Not*I and cloned into the vector PLEX (Invitrogen cat. No. K450-01) cut with *Bam*HI and *Not*I. The *E. coli* strain used was GI742 supplied by Invitrogen. The resulting plasmid insert (PLEX-*BgI*II/tn43, Figure 5) places the p14-2f and p13-1f genes under the control of the bacteriophage Lambda P<sub>L</sub> promoter. Figure 5 illustrates the cloning of DNA encoding nematocidal activity in the expression vector PLEX, where: A, plasmid clone; B, insert and gene locations; Tpr, trimethoprim resistance; Apr, ampicillin resistance; P<sub>L</sub>, bacteriophage lambda PL promoter; \*, plasmid joins to form a circular molecule; \*\*, incomplete genes. Selection of colonies on RMG media (described in the Invitrogen manual) containing ampicillin (50 µg ml-1) and trimethoprim (50 µg ml-1) prevents expression from the P<sub>L</sub> promoter. Colonies were then cultured on LB containing Trimethoprim (50µg ml-1) in 2.0 cm<sup>2</sup> wells for

nematocidal tests. The clone was active. This indicates that genes within this fragment have nematocidal activity. The clone PLEX-BglII/tn43 was digested with ClaI and religated, this resulted in a deletion of part of the p13-1f gene, this clone had reduced nematocidal activity indicating the importance of this gene.

All these results indicate that the genes and gene products of p13-1f and p14-2f are important for nematocidal activity. Other smaller genes within the *BgI*II to *Nru*I sites of cHRIM10 and PLEX-*BgI*II/tn43 may also be essential. In addition genes outside this region within the remaining cosmid clone (cHRIM5) may also encode products with nematocidal activity, or may enhance the nematocidal activity of genes in the smaller region (*BgI*II-*Nru*I of cHRIM10 and PLEX-*BgI*II/tn43).

### Example 8 - Field trials

Activity of strains selected in accordance with the above methods, or from depositary institutions which include bacteria which in nature are associated symbiotically with entomopathogenic nematodes, may be further assessed in field trials as follows.

Symbiotic bacteria in the absence of their nematode host can be inoculated into one or more portions of a field which is infested with nematodes, or into containers containing unsterilised soil from such a field. The bacteria can be inoculated onto the roots of plants, or into seeds. Periodically treated and untreated areas or containers can be assayed for nematode larva, egg, or cyst counts and for the presence of the inoculated bacteria by methods well known to those skilled in the art. A reduction in the number of nematode counts in areas in which the symbiote bacteria are present indicates control of the nematodes otherwise found in the untreated areas or samples.

#### Annex I - amino acid sequences

SEQ ID NO:1

P0-0f

ISWFATGIPTVDALLAEEFWHGDKQAFPPFTCRFTHFDPDKEQDVTLVPSTEEAYWLHRA
LQGQPLHSEVYGDDGTAQAGIPYTVMDSRPQVRLLTGLPGNSPTVWPSVIEQRTWQYERI
ADDPQCHQQVVLNSDRYGFPRETVDIAYPRRPKPAVSPYPDTLPATLFDSSYDEQQQQLR
LTRQRQHYHHLTDTEHQVLGLPDVMRSDAWGYPAARVPREGFTLEDLLAENSLIAPGTPL
TYLGHQRVAYTGTTGTEEKPTRQALVAYTETAVFDELALQAFNGTLSPEALEKKLIESGY
LSVPRPFNTGAESAVWVARQGYTDYGGSEAFYRPLAQRTTVQIGKNTLHWDTHYCAVVRM
QDAAGLYTDAAYDYRFLTPVQITDANDNQQHITLTALGQVSSGRFWGTEEGTPQGYTPPE
DRPFTPPSSVAEALDLKPDLPVANCMVYAPLSWMPLAHTYQEYIAGFTWQALLDAGVVTE
DKRVCALGFRRWVQRQGIVLNGQALADSREPVHVLTLATDRYDTDPDQQLRKSVTYSDGF
GRLLQSAVYHAPGEAWQRAADGSLITDAKGAPLVAHTATRWAVSGRTEYDGKGQPVRTYP
PFFLNAWQYLSDDSARQDLNADTHRYDPLGREYQVRTAKGYLRQNRLTPWFVVNEDENDT
LS

SEO ID NO:2

P1-2r

 ${\tt YLPQRGQCDMLLVVIGIGYLNGGQEAVIIGGIRVQTRRILHTDDRTVMGIPMEGVFANLH} \\ {\tt RRPLSQRTVKRLRPAVIGISLTGDPDRRFRTGIEWAWNRQITRLD}$ 

SEQ ID NO:3

#### P2-0f

SHLPARYGGRITTLSRKGFMTVNRGDNLHQKTPEVTVLDNRGLTVRELRYHRHPNTPTTT
DERITRHRFTLSGQLAHSIDPRLFDLQQTDNTVNPNMIYDTALTGEVVRTRSVDAGNDLI
LNDITGRPVLAINATEVTRTWQYENDTLPGRPLSITEQPAGEAGRITERFVWAGNSQAEK
NSNLAGQCVRHYDTAGLNQTDSIALNGIPLSVTRQLLPDGTDADWQGNNEPAWNDRLAPE
NFTTLSTADATGAVLTTTDAAGNLQRVAYDVAGLLTGSWLRLAGGTEQVIVKSLTYSAAG
QKLREEHGNGVVTTYTYEPETQRLVGIKTKRPQGHAQGTKVLQDLRYEYDPVGNVVKVTN
DAEVTRFWRNQKVVPENTYVYDSLYQLVSATGREMANIVQQSTLLPTPSLIDSSTYSNYS
RTYNYDRGDNLTQIRHSAPATGNSYTTDITVSDHSNRAVLDTLTDDPAKVDALFTAGGHQ
IPLQPGQNLVWTPRGELLKVAPVVRDGQISDQESYRYDAASQRIIKTHVQQTANSSQAQS
TLYLPGLERHTTINGTTVKEVLHVITIGEAGRAQVRVLHWENGKPGAISNNQMRYSYDNL
IGSSGLEVDGDGQIISMEEYYPYGGTAVWTARSQTEADYKTVRYSGKERDATGLYYYGYR
YYQPWAGSWLSADPAGTIDGLNLYRMVRNNPATLDDKNGLAPGNRYVFFPFIHEDRIFRL
ASANVYRTEHNKSDIIAVVEDKALDSKLFTNSIEQFFKKPKGKAILKGSPDIKERLLNNI
VHDLSNMQVGDQLYVNAHGHSAKPFFYSDSGYSKIIMEQLQRGANYVAKDLVNKFKLPEN
ATIKISTCHSAEGKGAHITVTSTGTNEKMRYSSIIENKGEFSRSLAGTMENELIKLQPGR

VRGNVYGYLGATTFYGAKNEKVIHLKDGNLTTGVHEGKLSMFTKKNRFSENIFGLKVKRS LTRTNFTGSGV

SEQ ID NO:4

p-2-9r

PAAEYVRDFTITCSVPPASRSQLPVSRPATSYATRCRLPAASVVVSTAPVASAVLRVVKF SGASRSFQAGSLFPCQSASVPSGSSWRVTDSGMPLSAILSVWFSPAVS

SEQ ID NO:5

P3-2r

QRALLNDIGHFAPGGTDQLIQAVIDIGVLRHHFLVAPEAGNLRIVRHFHHVPHRVVLIAQ VLQHLRPLCMSLWAFGFYANKALGLRLVGVGGHHAVAVLFAQFLTRGGIRQGFHDNLLCP ARKPQPTASQQACYVIRHTLQVTGRIGGGQYRAGGIRRAQGGEVFRCQPVVPGGFIVSLP VCVRTIRQQLARDGQRYAVKRNTVRLVQSGGVIVTHALSGQVAVLLRLTVPCPDKTLCDT ACFASRLFCDTERASG

SEQ ID NO:6

P3-6r

SDRRQTGYAYSADHYRISGRSTVCTVRAGLMNYQCWLQHAATQLSESDSPKRDAEILLGY VTGRSRTYLIAFDETLISSEELHQLDSLLVRRIQGEPVAYIIGEREFWSLPFAVSPATLI PRPDTECLVEKALELLPDSPARILDLGTGTGAIALALASERNDCYVTGVDINSDAVMLAQ HNAEKNAGKLAIHNVNFLQSEWFAAVGNQQFDMIVSNPPYIDERDPHLQEGDIRFEPATA LIAAQNGMADLQAIVGQARHFLSPNGWLLLEHGWKQGTVVRNLFLEKGYQQIATFQDYGG NERITIGRWNKNETHS

SEQ ID NO:7

P3-7f

SEQ ID NO:8

P5-6r

WQNGGSSSTTPRYLAGCYVWYPCSARLSGNAKSLLAPDGEWMKHTLKSKASGNTFTGRLI PTGRPTVVTIDKSGANTAALTLLNAEGEPQQGIEIRQNKYLNNRIEQDHRHVKRRIRPML GFKSFRRAQT

SEQ ID NO:9

WO 00/42855 PCT/GB00/00219

40

#### P5-7r

ALLFLSESRVMSLIRNAFKLLHYPVDIMAQCVRWSLTYALSLRNLEEMMAKRGIFVDHAT IPRWVLRLVPLLSKAFRKRKKPVGSRWRMDETYIKVKGQWKYLYRSVDTDGQTDCGDYR

SEO ID NO:10

#### P6-3f

 $\label{thm:continuous} \begin{tabular} $\mathsf{VHSPSGAVAPGKFFIENFADTFPAPLPLHPFIDACIQQGFQLLPCLIAIAHSGKQAFECV$$ $\mathsf{LLDRLALQGSQCLQALVLPVGDVNGQTAHGFLLIGYTQTHISTYNGLWLFITQGVRYRFV$$ $\mathsf{RQTFVCRSLSFSEDDCTN}$$ \end{tabular}$ 

SEQ ID NO:11

#### P6-3r

RTCRERPRLMDYVLTKAAEADLRAIIRHTRKQWGDAQVRRYITALEQGIARLAVGQGSFK DMSALFPALRMAHCERHYVFCLPRENAPALIVAIFHERMDLLTRLADRLK

SEQ ID NO:12

#### P6-6r

PQTIICANVGLCITDKEKTMSRLTIDITDRQHQSLKALAALQGKTIKQYALERLFPGMSD SDQAWQELKALLDTRINEGMEGKGCGKSIGEILDEELAGSDRA

SEQ ID NO:13

#### P7-1f

NAHFLIVSKTNVVMSNQDPHNKRDSLFSAPIANLGDWSFDERVAEVFPDMVKRSIPGYSN IISMIGMLASRFVTPGSQIYDLGCSLGAATLSIRRSINADNCRIIAIDNSPAMIERCRRH IDSFKASTPVEVIEQNILDTDIQNASMVVLNFTLQFLHPDDRQKILKKIYAGLKPGGVLV LSEKFNFEDQKIGELLFNMHHDFKRANGYSELEVSQKRSMLENVMRTDSVDTHKSRLKEV GFQHVEVWFQCFNFGSLLAIKGTEQ

SEQ ID NO:14

#### P7-9f

TMIDFGNFYQLIAKHPLNHWLDSLPAQLSHWQKTSQHGQFSSWVKILENLPEIKPSHLDL KNGVIAIHEPDLSKGEKARLHNILKILMPWRKGPFSLYDVEIDTEWRSDWKWERVLPHIS PLEGKTVLDVGCGSGYHMWRMVGEGAQLVVGIDPTQLFLCQFEAIRKLLGNNQRAHLLPL GIEQLPELQAFDTVFSMGVLYHRRSPLDHLWQLKNQLVSDGELVLESLVIEGDENQCLIP GERYAQMRNVYFIPSAKMLKVWLEKCGFVDVRIVDHAATTPDEQRRTEWMKTESLVDFLD PSDHSKTIEGYPAPLRAVLIARKP

SEQ ID NO:15

#### P8-4r

SLQIDREKVGLDRYPQPIERLRQPCATCDNHCHSRHQVRFFLLKEKYGAALAPISSQSAI RYQFQRHTMKKGLFAMASIFSGYCGGELFHLLTDPAHESQ

SEQ ID NO:16

#### P9-8r

SSFRLNDDLLTNSYSEGFLMIKLEICCYSISCALVAQNAGADRIELSASPLEGGLTPSFG
ALQQSLQRLSIPVHPIVRPRGGDFCYNNMDFEAMKNDVARIRDMGFPGIVFGILSENGHI
DRLRMRQLMSLSGNMAVTFHRAFDMCFNPHVALEQLTELGVQRILTSGQQQNAELGLTLL
KELMQASRGPIIMPGAGVRVSNISKFLEAGMTEVHSSAGKIVPSTMKYRKVGVAMSSDDR
DVDEYSHYSVDGELVESMKGVMSLIKR

SEQ ID NO:17

#### P10-5r

YFGKNRRFVIYVTLMERNFYGLFNGEEMSHFSKISELQDLVADLAGFEQKLKQFEGHLGL HFEQYSADHISLRCNESKIADRWRKGFLQCGQLISESIINGRPICLFDLNQPIVLLDWKI DCVELPYPSQKHYVHQGWEHVELVLPVPPEQLICEAKKLLPQPLPDNFRMKESHPKGKNE RLPNPILAV

SEQ ID NO:18

#### P10-7f

GNTVNIQVILSEKISNALIEAGAPTDSEAHVRQSAKAQFGDYQANGVMAAAKKVGIPPRQ
LAEKVVSQLDLQGIASKVEIAGPGFINIFLDKAWVAANIETTLKDEKLGITPVEPQTIVI
DYSAPNVAKQMHVGHLRSTIIGDAAARTLEFLGHKVIRANHVGDWGTQFGMLIAYLEKIQ
NENANDMALADLEAFYREAKKHYDEDEEFAIRARNYVVKLQGGDEYCRKMWRKLVDITMS
QNQETYNRLNVTLTEKDVMGESLYNDMLPGIVADLKQRGIAVKSDGATVVYLDEFKNKEG
EPMGVIIQKKDGGYLYTTTDIACAKYRHETLNASRVLYYIDSRQHQHLMQAWAIVRKTGY
IPESMSLEHHMFGMMLGKDGKPFKTRAGGTVRLSDLLDEAIERADTLIREKNPDMPEDEL
KKVVEAVGIGAVKYADLSKSRTTDYVFDWDNMLAFEGNTAPYMQYAYTRVSSIFKRADID
ENSLTLPVMLNEEREQALATRLLQFEETITTVAREGTPHVMCAYLYDLAGLFSGFYEHCP
ILNADSEELRQSRLKLALLTAKTLKQGLDTLGIQTVERM

SEQ ID NO:19

#### P11-1r

AQVSNMHLLGDIRCGIIDNDGLRFHWGDTELFIFQGSFYICCNPRFIKKNIDKTWACNFN FAGNSLQIQLADDFFCQLSRRYSHLFSGSHHTIRLIVTKLCFGRLTDVSFTVGWSASFNQ RIADFF WO 00/42855

42

SEQ ID NO:20

P12-1r

HARVGVLHIRCRVAFKGQHIIPVENIVCSTALGKICIFHRANPYRFHDFFQFVFWHIWVF LTNEGIRTLNRFIQQIGQSYCAAGTGFEWFTIFAQHHAKHVVFE

SEO ID NO:21

P12-5r

YHASFQLCRRLLHTFYSLNTQSIKTLLQSFRCQQSQLQAALAQFFAIGIQDRAVLIETRE QTGQIVQVCTHNMWRTFTGDGSDRFFKLQQAGCQCLLAFFIQHHRQCQAVFIDIRTFKDR

SEQ ID NO:22

P13-1f

FTLREDSMSDWTGVSTFNVILETGLDNCNIYANGLNMIGVIINITPTDDEGNFVDIDDVT LNDNIKIVDYIDGSDIDGSDGWFYTGNPNEYNTIPNSQSYSLLKSENSQITQIKRYVSCS NTSRLRTKSFSAKVTTTSGKVISITQNSINSSRVVINAIDATNFTDDELRTTKETRFENQ SYTSHKSSTNSLYVHTWTIPRSLKLQNWRWEDYNNGWTWAQSCYYKTGADGGSESTRWLA AGSIFPPGNYDGLWLDNDIALSGMAHKSYNVDTGINQLSFTRIIGKGFSWVYNISGLDRG HAVIIIDQYGNKYRILFHAGYENSDPYLSSSIVY

SEQ ID NO:23

#### P14-2f

VYIKFLKLFRRITMSDNNEFFTQANNFTSAVSGGVDPRTGLYNIQITLGHIVGNGNLGPT LPLTLSYSPLNKTDIGFGIGFNFGLSVYDRKNSLLSLSTGENYKVIETDKTVKLQQKKLD NLRFEKDLKENCYRIIHKSGDIEVLTGFNNNAFDLKVPKKLLNPAGHAIYIDWNFEATQP RLNRIYDDLDGHDIPLLNLEYQGLIKTILTLFPGQKEGYRTELRFLNRQLNSIHNFSLGN ENPLTWSFGYTPIGKNGILGQWITSMTAPGGLKETVNYSNNNQGHHFPQSANLPVLPYVT LMKQVPGAGQPAIQAEYSYTSHNYVGGGSNGIWNNKLDNLYGLMTEYNYGSTESRRYKDK EGHDQIVRIERTYNNYHLLTSECKQQNGYIQTTETAYYAIIGHNFDSQPSQFQLPKTKTE TWRSADNSYRSEITETTFDESGNPLTKVIKDKKTQKIISPSTHWEYYPPAGEVDNCPPEP YGFTRFVKKIIQTPYDSEFKDDPEKFIQYRYSLIGSQSHVTLKIEERHYSATQLLNSTLF QYNTDKSELGRLLKQTECTKGENGKTYSVVHKFTYTKQDDTLQQSHSITTHDNFTIHRSQ VRSRYTGRLFSDTDTKDIVTQMSYDKLGRLLTRTLNSGTPYANTLTYDYELNNLQDDNRP PFVITTTDVNGNQLRNEFDGAGRHVSQCLKDSDGDGKFYTIHTQQYDEQGRHHTSTYSDY LTNGRQQTDPDKVHLSMSKSYDNWGQIANTHWSYGVSEKITVDPITLTATKQLQSNSNNV QTGKEVTTYTPSQQPIQITLFDEAGHLQSCHTLTRDGWDRVRKETDAIGQCTIYQYDNYN RVIQITLPDGTIVNRKYAPFSTDTLITDIRVNGISLGQQTFDGLSRLTQSQDGGRVWAYT YSAGNDQCPSTVITPDGQFIHYQYQPELDDAVLQVASNEITQQFSYNPVTGALLKAVAEG QSLTPIYYPSGRLKMENINDMKKMSYLWTLRGLENGYTDLTGTIQKISRDTHGRVTQIKD

SSIKTTLNYDDLNRHIGSQVTDLATGHMLTTTVEFDGLNREIGRKLCDSSGHTLDIQQSW
LKTQQLANRIVKLNGVLQRTEQYSYDSRNRLNQYKCDGAECPTDKYGHSIVTQNFTYDIY
GNITACHTTFADGTEDHATFKFANPTDPCQLTEVHHTHPDMPDNIRLKYDKAGRVINITD
NHGNTENFTYDTLGRLQNGQGSVYGYDPLNRLVSQKTDTLDCELYYRETMLVNEVRNGEM
IRLLRTGETIIAQQRASKVLLTGTDSQQSVILTSDKQNLSQEAYSAYGKHKSTANDASIL
GYNGERADPVSGVTHLGNGYRSYDPTLMRFHTPDSLSPFGAGGINPYSYCLGDPINRSDP
SGHLSWQAWTGIGMGIAGLLLTIATGGMAIAAAGGIAAAIASTSTTALAFGALSVTSDIT
SIVSGALEDASPKASSILGWVSMGMGAAGLAESAIKGGTKLATHLGAFAEDGENALLKST
SESSRIKWGVTRSLDREIVRNEEGQVIKDHSRGYTDNFMGKGEQAILVHGDKDGFLYHTE
GNKHNGKGPYTRHTPEQLVDYLKDNNIVDLTQGGDKPVHLLSCYGKSSGAADKMAKYINR
PVIAYSNKPTISQGLARIERKDFFLKSTYHSYDPRKIILGRTEKTVKPKTFRP

SEQ ID NO:24

P17-6r

LCYGHICLSGIPHRHIYIGSTYYGNRKSTVLYAAILHSVSLFYLLIAVFSASSAGYLTYG LSYHTISVQFLGLSHQIPLLLSTYDQSLNLLLDYQYGDSGHRNLE

SEQ ID NO:25

P17-8r

SAQCIVGKVFRISMVISDIYYSTSLIIFQPDIIRHIWMSVVYLCQLAWVSWVGKFEGSMV FCPICECGVTGGDIAIDIISKILCDYAMAIFVCRAFRTVTFILVQPITGIVRVLFCTLQY SIQFHYSIC

SEQ ID NO:26

P18-7r

PSSLRTISLSKLLVTPHFILELSEVDLSKAFSPSSANAPRCVASLVPPLMADSANPAAPI PIETHPSIEDAFGEASSSAPLTIDVISDVTLSAPNASAVVEVEAIAAAIPPAAAIAIPPV AMVSSNPAIPMPIPVHACOLK

SEQ ID NO:27

P19-5f

AHCHIALFPCWHNPQYCQQHPDHHSNCHHQFKQEYPPSRQRRENITLTQLPIKHTGIEAG SQTNRKRQTCMFQRANESKVHOLGONOGRDRNFYWCFDILT

SEQ ID NO:28

P19-8f

PQSTPSSQNSRQLTPAESSQHQKQKSDHIEIMIPSEAPREYREQLHKATPARNRDVAPNP SVFDILRDYHWKNFSPVKAAKSSLTPHPVHQKAIPLNDQRNTSMKQSLKPEMRQKLY

SEO ID NO:29

P20-1r

 $\label{topolicy} {\tt GKNCINDQGNLPDRYTQNCRPHLTDNPPYGTVTERNPRQYQHADLFQMRKLIGQLQNPSG} \\ {\tt NNGPTQRQHWRIAIRSHKQCKNDHTDIEQCRSKSRHRKAVPCIKNCASQRSQRNQKDIRK} \\ {\tt RNSK} \\$ 

SEQ ID NO:30

P20-9r

NNTMNLLKSLAAVSSMTMFSRVLGFIRDAIIARIFGAGMATDAFFVAFKLPNLLRRIFAE
GAFSQAFVPILAEYKNQQGDEATRTFIAYISGMLTLILAIVSVIGVIAAPWIIYVTAPGF
TDTPDKFVLTRDLLRITFPYIFLISLASLAGAILNTWNRFSVPAFAPTLLNVSMIIFALF
VAPYCNPPVLALGWAVVAGGVLQLAYQLPHLKKIGMLVLPRISFRDSAVWRVIRQMGPAI
LGVSVGQISLIINTIFASFLVSGSVSWMYYADRLMELPSGVLGVALGTILLPSLAKSFSS
GNHEEYRKLMDWGLRLCFLLALPCAVALGILAEPLTVSLFQYGHFSAFDAEMTQRALIAY
CFGLMGLIVVKVLAPGFYSRQDIKTPVKIAIATLILTQLMNLAFVGPLKHAGLALSIGLA
ACFNASMLYWQLRKRDIFTPLAGWGIFLFKLVVAIAVMVGVLLAVLWVMPAWEQGNMAMR
LLRLMGVVIAGAGSYFAVLALMGFRLKDFAHRGLQ

SEQ ID NO:31

P21-7r

AIILIRDKLSRIFSRQISGEGMFGYRSASPKIRFITDRMVVRLVYERDAYRLAEYYSENK DFLKPWEPTRDGSFYQPSGWTNRLNYIAELQRQNATFNFVLLDSDEREIMGVANFTNVVR GAFHSCYLGYSLAEKLQGQGLMYEALQPAIRYMQRYQRMHRIMANYMPHNHRSGNLLKKL GFEQEGYAKNYLMIDGVWQDHVLTALTDDAWGKVGL

SEQ ID NO:32

P21-8f

WCAMSLVSQARSLGKYFLLFDNLLVVLGFFVVFPLISIRFVEQLGWAALIVGFALGLRQL
VQQGLGIFGGAIADRFGAKPMIVTGMLLRALGFALMAMAHEPWILLLSCVLSGLGGTLFD
PPRAALVIKLTRPHERGRFYSILMMQDSAGAVVGALIGSWLLQYDFNIVCWIGASIFVLA
ALFNAWLLPAYRISTIRTPIKEGMMRVIRDRRFLYYVLTLTGYFVLSVQVMLMFPIIIHE
ITGTPTAVKWMYAIETAISLTLLYPIARWSEKHFRLEQRLMAGLFLMSICMFPIGWVNQL
HTLFGLLCLFYLGLVTADPARETLSASLSDPRARGSYMGFSRLGLALGGAIGYTGGGWLY
DTGRDLNMPQLPWILLGLSGLITIYALHRQFNQKKIDPVMLGRH

SEQ ID NO:33

P23-1f

KGANMKRFFLGAALVLVGLVSGCDQFKDFSINEGLMNDYLLKKVHYQKKISIPGIANANI TLGDLSSQIGRQDPEKIELSTQAKVQLATLLGTIQADMKLTIKAKPVFDAEKGAIFVKGL EIVDYQTTPEKAAAPVKALIPYLNTSLSEFFDTHPVYVLNPEKSKAEAAASQFAKRLEIK PGKLVIGLTDK

SEO ID NO:34

P24-4r

QVALQHGRRLGTITLFDNLLGLNQVMNEFSIVCRILGTLFNRAPQDPVLQPLITMIAEGK LKQAWPLEQDEWLDRLQQNSELSVMAADYHALFTGESASVAVCRSDYTDGEESEVRQFLT ERGMPLSDTPADQFGSLLLAVSWLEDQAAEDEIQAQITLFDEYLLPWCGQFLGKVEAHAT SGFYRTLAIVTREALQALRDELESE

SEQ ID NO:35

P25-3r

DCMNIIFFHPSFNTDEWIQGIQARLPDAKVRQWVSGDQEPADYALVWQPPYEMLANRQGL KGIFALGAGVDAIFKQESKNPGTLLADVPLIRLEDTGMGRQMQEYAITSVLHYFRRMDEY KRYQEQRLWNPIAPHNRKEFVIGVLGAGILGRSVIGKLMEFDFNVRCWSRTSKQLDSVES FYGKEQLGDFLSGCKVLINLLPDTPDTRGILNLSLFSQLKSGSYVINLARGAQLVEQDLL VAIDKGYIAGATLDVFAEEPLSNMHPFWTHPRINVTPHIAANTIPEAAMDVICENIRRMV QGEMPTGLVDRVRGY

SEQ ID NO:36

P26-0f

 $\tt KTSQGFTSTTCSNGNVLKICGLITPCSSLIQRTYPNNMTIGIFSKESTAKNFGMGFLYYF\\ DLRVLSPFFKAPINIFTGWQHNTNFRKSRNSTIRLCSSTPNSKQYFTTSRKCHITGAGKY\\ RFSIENCFIKSG$ 

SEQ ID NO:37

P27-0r

YSAGCSTVLKSSLNLQCDTFNCESFVMLTLNFSTSVNAKPSHIWAHYVDFDLRKKWEVDL EYFQFEGEVKTGQYGRMILSGMPEIRFYLSNIEVNKEFTDQVNLPQMGILTFRHQIITDE NNMACRVQVTVSFEPDANIPAVQAESFFKQGTQDLVESVLRLKSVVETVSPKPNLQLVYV SDIESSTAFYKTIFNAEPIFASSRYVAFPAGGEVLFAIWSGGAKPDRAIPRFSEIGIMLP SGKDVDRCFEEWRKNPEIKIVQEPHTEVFGRTFLAEDPDGHIIRVCPLD

SEQ ID NO:38

P27-8r

KGNQITMILYKGSKNYLFNQLNYDSCVLLEVDESVNLNGWDELSRAQRLLFLMEILRRYH

WO 00/42855 PCT/GB00/00219

46

FPVQGKVLAQKLNISLRTLYRDIASLQAQGAIIEGEPGIGYVLRPGFVLPPLMFTQNEIE ALALGANWVAKRADPQLKESANNAISKIAAVIPAELKQMLEASSLLIGPAATAVQPVVEI QQIRQAINTRHKITLAYLDIKDIPSERTIWPFALGYFENISIVIGWCELREEFRHFRSDR IMRLKIENQCYPRSRQVLLKEWRAMEKISR

SEQ ID NO:39

P27-9f

RKMTIYDLKPRFQNLLRPIVIYLYKQGITANQVTLTALFLSIFAGSLLSLFPSPHLYWLL PVFLFIRMALNAIDGMLAREHNQKSHLGAIYNELGDVISDVALYLPFCLLPDVNSLSLLI ILFLTILTEFIGVLAQTIGASRRYDGPIGKSDRAFIFGAYGLIIAIFPLALGWSISLFAF MIILLLVTCYQRVVKALREIRLAEOSHSK

SEQ ID NO:40

P28-5f

GVNMTPQLDQRIAEEHYFTTSDNASLFYRYWPQQQANPDRAIIIFHRGHEHSGRIQHVVD GLDLPDVPMFAWDARGHGKTEGPRGYSPSMGTSIRDVDEFVRFIATQYGIAMENIVVIGQ SVGAVLVSAWVHDYAPKIRAMILAAPAFDIKLYIPFATQGLQLMQKARGIFFVNSYVKAR YLTHDETRIASYNSDPLITREIAVNILLDLYQTAERVVKDAAAITLPTLLFISGSDYVVN KKPQHQFYQQLNTPIKEKHVMDGFYHDTLGEKDRHLVFDKIRVFIERIFALPRYQHDYSQ EDTWSHSADEFRTLSTSLPCLCPKKLSYQLMRKVMSTHWGRTSEGVCIGLKTGFDSGSTL DYVYRNQPQGKGILGRILDKHYLNSIGWRGIRQRKIHIEMLIRHAIRSLREQNMPVHMVD IAAGHGRYILDAINDFSKVDSILLRDYSEINVNQGQAYIEERDLTDKIRFIIGDAFNAES ISSITPAPTLGIVSGLYELFPDNNLLRNSLRGFADVMTENGYLVYTGQPWHPQIEVIARV LSSHRDSQPWIMRRRTQGEMDALVEAAGFEKLYQLTDNWGIFTVSIAKRVHR

SEQ ID NO:41

P28-5bf

HHNSINVLLKNIISPHQIMLLCFTVTGHNNRPIQTERSLFFTVVMSTQDVSSMSLTDSIC LMFLCSRGMPVDTVRQKGRAVTAHPWERRFVMLMNLSDLLPLSTASPWKISWLSARVSER Y

SEQ ID NO:42

P30-3f

INKYKMEHHMHSSLDSRRRLWLTGVIWLLFLAPFFFLTYGQVNQFTAQRSDVGTVMFGWE HNIPFWSWSIIPYWSIDLFYGISLFICTHRREQWLHGWRLMTASLIACVGFLLFPLKFSF SRPTTEGLFGWLFNQLELFDLPYNQAPSLHIILLWLLWLRYSAYVSGYWRGLLHIWSVLI ALSVLTTWQHHFIDVLTGFAVGVILSYLLPVSYRWRWQPNQDRYARKLFGYYLTGSALFA LIASLLGGSFWILLWPAVSLLMIALGYAGLGSSVFQKQPDGRMSLSARWLLAPYQLGAWL SYLWFRRKSAPFNHITEGIILGSLPCQPVTAVSVLDITAEWHRRSDARTVNYVCOPOIDL WO 00/42855

47

SEQ ID NO:43

P31-6f

QSCVKPDRMSRSDKHIWMPCLNGQKATYNGEHNMQPENLISKVIIATLKSWRFISTLSAF SILIATAMLIAVFNTTALNNIALYAVLLFTTLYCQYYCWRTWLDCHYFQILNSSPEKSAE FDQTLLLIFNKLPQSRTQNDRFNGAIKLLKKATIGLILQWILFFLFLLTLKYSA

SEQ ID NO:44

P32-3f

MNTRKINGIRPFSAFIDSCLKESYSFPRFIRDIIAGITVGVIAIPLAMALAIGSGVAPQY
GLYTAAIAGIVIAMTGGSRYSVSGPTAAFVVILYPVSQQFGLSGLLIATLMSGVILIVMG
LARFGRLIEYIPMSVTLGFTSGIAITIATMQVQNFFGLKLAHIPENYIDKVVALYQALPS
LQLSDTLIGLTTLLVLIFWPKLGVKLPGHLPALIAGTAVMGAMHLLNHDVATIGSSFSYT
LADGTQGQGIPPILPQFVLPWNLPDTHSLDISWNTVSALLPAAFSMAMLGAIESLLCAVI
LDGMTGKKHHSNGELLGQGLGNIAAPFFGGITATAAIARSAANVRAGATSPIAAVVHSLL
VLLTLLVLAPMLSYLPLAAMSAILLIVAWNMSEAHKVVDLIRHAPKDDIIVMLLCLSLTV
LFDMVRRDHYRHCAGITPVYAQNCQYDSNQHVIFNKRGERVIGRTN

SEQ ID NO:45

P33-4r

ESIGAKTSNVNNTSRECTTAAIGEVAPARTLAAERAIAAVAVMPPKKGAAILPNPWPSSS PLEWCFFPVIPSRITAHSNDSIAPSMAIENAAGSNADTVFQLISRECVSGKFHGRTNWGR MGGMP

SEQ ID NO:46

P33-5f

LSYSIWSVAITIGIVLASLLFMRKIANMTRISTSSLTSAEKGLLVVRINGPLFFAAAERI FAELREKSADYQTIIMQWDAVPVLDAGGLHAFQGFVRELGKEKHIVVCDIPFQPLKTLAR AKVMPIEGELSFYATLPKALKEMAVDYTPEVCASSEKIQGQ

SEQ ID NO:47

P34-3f

CMSDVENDRRTLGSLLHDTEAQHVNHQIVITKVAATVTQDHLVIAAFFEFFNNIAHLPRA NKLWFFNINHSTGFRHRFNQIGLAGKEGWKLNHIHHIRDWLSLCRLMHVSDNFHAEGLFQ FLKDFHPLFQPWPTIRADRRTVSLIKRRFKNIRNAQFLCHGDIVLTNPHGQIP

SEQ ID NO:48

#### P35-0r

LSCIRFIFLLIQQIYLPLTREGISMQQKVVNIGDIKVANDLPFVLFGGMNVLESRDLAMR ICEHYVTVTQKLGIPYVFKASFDKANRSSIRSYRGPGLEEGMKIFQELKQTFGVKIITDV HEPAQAQPVADVVDVIQLPAFLARQTDLVEAMAKTGAVINVKKPQFVSPGQMGNIVEKFK EGGNDQVILCDRGSNFGYDNLVVDMLGFGVMQQATQGAPVIFDVTHALQCRDPLGAASGG RRAQVAELARAGMAVGIAGLFLEAHPDPENAKCDGPSALPLAKLESFLMQIKAIDDVVKN FPELDTSK

SEO ID NO:49

#### P35-8r

VDGIKMKPIVNYEFNNTPLIDGIILVSKIIRPDFPQTLVSEQLTALVEEARQRLSSITDS
KVKLDSLLTLFYREWKFGGANGVYCLSDTLWLDRLLHSRQGSPVSLGTVFTHIAQALGLS
VQPVIFPIQLILRIDLLDQPTWFINPLNGDTLNEHTLDVWLKGNIGPTVRLKKQDLQEAD
NVSLVRKITDTIKVSLMEEKKMELALKASEVVLTFDPDDPYEIRDRGLIYAQLDCNHIAV
SDLSYFVEHCPEDPISEMIKMQINTIEQRLIVLH

SEQ ID NO:50

#### P36-7r

SDRRQTGYAYSADHYRISGRSTVCTVRAGLMNYQCWLQHAATQLSESDSPKRDAEILLGY VTGRSRTYLIAFDETLISSEELHQLDSLLVRRIQGEPVAYIIGEREFWSLPFAVSPATLI PRPDTECLVEKALELLPDSPARILDLGTGTGAIALALASERNDCYVTGVDINSDAVMLAQ HNAEKNAGKLAIHNVNFLQSEWFAAVGNQQFDMIVSNPPYIDERDPHLQEGDIRFEPATA LIAAQNGMADLQAIVGQARHFLSPNGWLLLEHGWKQGTVVRNLFLEKGYQQIATFQDYGG NERITIGRWNKNETHS

SEQ ID NO:51

#### P37-5r

VEMREMAQEELKEAKIRNEELEQQLQLLLLPKDPDDERNCFLEVRAGTGGDEAAIFAGDL FRMYSRYAEARRWRVEIISANEGEHGGYKEVIAKVSGDQVYGHLKFESGGHRVQRVPETE SQGRIHTSACTVAVMPEIPEAELPDISPGDLKIDTFRSSGAGGQHVNTTDSAIRITHLPT GIVVECQDERSQHKNKAKAMSVLAARIRAAEMRKRQEVEASERRNLLGSGDRSDRNRTYN FPQGRVTDHRINLTLYRLDEVIEGKLDMLIQPIIIEYQADOLSALSEOD

#### Claims:

- 1. The use of a bacterial strain to control a target nematode, characterised in that in nature the bacterial strain is associated symbiotically with an entomopathogenic nematode.
- 2. The use according to claim 1, wherein the bacterial strain from nature is directly employed to control the nematode target, or is employed to give a recombinant bacterium employed to control the nematode target, or the natural or recombinant strain is employed as a source of a nematode control agent to control the nematode target.
- 3. The use according to claim 1 or 2, wherein the target nematode is not the same as the nematode with which the bacterial strain is found symbiotically in nature.
- 4. The use according to claim 1, 2 or 3, for control of helminthiasis in a human or a domesticated animal or the control of plant pathogen nematodes.
- 5. The use according to any preceding claim wherein the nematode to be controlled comprises one or more of Haemonchus, Trichostrongylus, Ostertagia, Nematodirus, Cooperia, Ascaris, Bunostomum, Oesophagosromuni, Chaberria, Trichuris, Strongylus, Trichonema, Dictyocaulus, Capillaria, Heterkis, Toxocara, Ascaridia, Oxyuris, Ancylostoma, Uncinaria, Toxascaris, Parascaris, Aphelenochoides, Anguina, Bursaphalenchus, Criconemella, Melodigyne, Ditylenchus, Globodera, Heliocotylenchus, Heterodera, Pratylenchus, Radopholus, Rotelynchus, Tylenchus, Trichodorus, Xiphenema, and Caenorhabditis.

- 6. A composition for the control of parasitic nematodes which comprises as an effective agent a species of bacterium which is a symbiont of an entomopathogenic nematode, or an engineered bacterium, or a nematode control agent derived from a natural or engineered bacterium.
- 7. A composition according to claim 6, wherein the bacterial species is of the genera Xenorhabdus or Photorhabdus,
- 8. A composition according to claim 7, wherein the bacterial species is of the genus *Xenorhabdus*
- 9. A composition according to claim 8, wherein the bacterial species is of, the species *Xenorhabdus bovienii*.
- 10. A composition according to claim 8, wherein the bacterial species is: Xenorhabdus bovienii strain H31 deposited with NCIMB under accession number NCIMB 40985; Xenorhabdus bovienii strain I73 deposited with NCIMB under accession number NCIMB 40986; and Xenorhabdus strain C42 deposited with NCIMB under accession number NCIMB 41004.
- 11. A composition according to any of claim 6, wherein the nematode control agent which is derived from a symbiont of an entomopathogenic nematode or from an engineered bacterium has functional activity against a nematode, and is a peptide.
- 12. A nucleic acid encoding a peptide of claim 11.
- 13. A nucleic acid according to claim 12, which nucleic acid comprises a

natural nucleotide sequence or a degeneratively equivalent sequence, or a functional variant thereof.

- 14. A nucleic acid according to claim 13, which is a homologous variant encoding a peptide which is a nematode control agent, the nucleic acid having 70% or more DNA sequence identity and/or the peptide having 70% or more amino acid sequence identity.
- 15. A nucleic acid according to claim 13, which is all or part of cosmid cHRIM5, in particular p 13-1f or p 14-2f, and variants thereof.
- 16. A nucleic acid according to claim 13, 14 or 15, wherein the variant has a sequence which is a derivative by way of addition, insertion, deletion or substitution of one or more nucleotides.
- 17. A nucleic acid according to any of claims 12 to 16, which is part of a longer sequence and the nematode control agent is expressed as a fusion protein.
- 18. A nucleic acid complementary to a nucleic acid according to any of claims 12 to 17.
- 19. A nucleic acid for use as a probe or primer having a nucleotide sequence of at least 15 nucleotides, which sequence is present in a nucleic acid according to any of claims 12 to 18.
- 20. A method for identifying or cloning a nucleic acid according to any of claim 12 for a nematode control agent, which method employs a nucleic acid probe according to claim 19.

PCT/GB00/00219 WO 00/42855

52

- 21. A method according to claim 20, which comprises the steps of:
  - providing a preparation of nucleic acid from a bacterium, (a)
  - (b) providing a probe,
  - contacting nucleic acid in said preparation with said probe (c) under conditions for hybridisation of probe to any said gene or homologue in said preparation, and,
  - (d) identifying said gene or homologue if present by its hybridisation with said probe.
- 22. A method according to claim 20, which comprises the use of two primers to amplify a nucleic acid encoding a nematode control agent, at least one of the primers having a conserved nucleotide sequence of at least 15 nucleotides.
- 23. A method according to claim 20, which comprising the steps of:
  - providing a preparation of nucleic acid from a bacterium, (a)
  - (b) providing a pair of nucleic acid molecule primers, at least one of which is a primer,
  - contacting nucleic acid in said preparation with said primers (c) under conditions for performance of PCR,
  - performing PCR and determining the presence of absence of an amplified PCR product.
- 24. A recombinant vector comprising a nucleic acid according to any of claims 12 to 17.
- 25. A host cell containing a vector according to claim 24 capable of replication.
- 26. A host cell according to claim 25 which is a plant cell.

- 27. A method for producing a transgenic plant which comprises the step of regenerating a plant from a plant cell according to claim 26.
- 28. A plant produced according to claim 27 which is a crop species which can be maize, cotton, soya, rice, *Brassica* species, tomato, potato, sugar beet, barley, soybean, peanut, onion, rye, wheat, corn, banana, raspberry, bean, or a decorative or other plant.
- 29. A method of producing a peptide nematode control agent comprising causing or allowing expression of a nucleic acid according to claim 12.
- 30. An antibody or fragment thereof, or a polypeptide comprising the antigen-binding domain of the antibody, capable of specifically binding a peptide of claim 11.

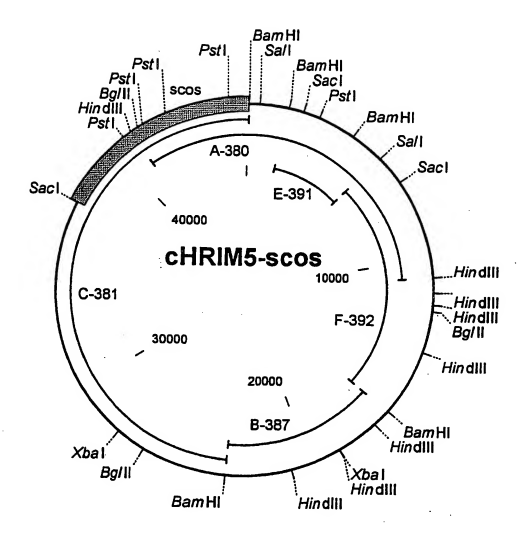


Fig. 1

2/14

			2/17			
Molecule:			Sequence I			
Description	: chrim	n5ed2.seg,	37544 bps I	ONA ·		
1 g	gatcagctg	gtttgccacc	gggatcccca	ccgttgatgc	cctgttagcg	gaggaattct
61 g	gcacggtga	caaacaggct	ttcccgccct	ttacctgccg	ttttacgcat	tttgaccctg
121 a	taaagaaca	ggatgttact	ctcgttccct	cgacggaaga	ggcttattgg	ctgcaccggg
181 c	gttgcaagg	ccaaccgtta	cacagtgagg	tctatggcga	cgatggcacc	gcgcaggcgg
	tarececta	taccgttatg	gacagtcggc	cccaggttcg	gcttctgacg	ggtttaccgg
361 t	taccastas	tocageetgg	ccgagtgtga	ttgaacagag	aacctggcag	tacgaacgga
421	аспопапас	catcancatt	acttatora	cggcgccgaa	cagtgaccgc gcctgcggtg	tacggttttc
481	ggatacgct	accadeasea	ttattcgaca	gccyccctata	tgagcagcaa	caccattac
	gcttacccg	gcaacggcaa	cattaccatc	acctgactga	cactgaacat	caactactaa
	actgcctga	tgtcatgcga	agcgatgcct	ggggctatcc	ggcagcgcgg	gtaccccgtg
661 a	aggtttcac	cctggaggac	ttgctggcag	agaacagtct	gatageceeg	ggcacgccat
721 t	gacctattt	agggcatcaa	cgcgtggctt	ataccggaac	gaccggaacc	gaagaaaaac
781 c	gacccgaca	ggcgctggtg	gcttataccg	aaaccgcggt	ttttgatgaa	ttagecttac
841 a	ggcctttaa	tggcacattg	agtcctgaag	ccctggaaaa	gaaattaatc	gagtctggtt
901 a	tttgtctgt	tccacgccca	ttcaataccg	gtgcggaatc	ggcggtctgg	gtcgcccgtc
961 a 1021 c	gggatatac	cgattacggc	gggtctgagg	cgttttaccg	tccgttggct	cagcggacga
	acaaastac	agcagateta	tacaccatt	gggataccca	ttactgtgcg ttaccgcttc	gtcgtccgta
	tcagataac	coatoccaat	dacaaccadc	aacatatcac	actgaccgcg	ctgaececeg
	atcatccgg	ccaattctaa	ggcactgagg	aagggactcc	gcagggttat	accedecte
1261 a	agaccgccc	atttacgcca	ccgtcctcag	tggcggaagc	cctcgacttg	aaaccogatc
1321 t	tccggttgc	caactgcatg	gtttatgcgc	cactgagtta	gatgccgttg	gcgcacacct
1381 a	tcaggaata	tatagccggc	tttacgtggc	aggcactgct	tgacgcgggg	gtagtgacgg
1441 a	agataagcg	ggtttgtgcg	ctgggtttcc	gtcgctgggt	gcaacgtcag	ggcattgtgc
1501 to 1561 as	gaatgggca	ggcattggcc	gattcacggg	aacccgtcca	tgtcctaacg	ctggccactg
	ccgttatga	cacggatece	gatcagcaac	tgcgcaagag	cgtcacctac	agcgacggct
	agatggcag	cctgatcacg	geageetace	acgegeeagg	agaagcctgg cgtagcccat	caacgcgcgg
	ctaaacaat	ctcaggcagg	acadagtato	accotaaacc	gcaacccgtc	casacctacc
	gccattctt	cctgaatgcc	togcagtacc	tcagtgatga	cagtgcacgg	caggatttaa
1861 a	tgccgatac	acaccgttat	gacccgctcg	gccgggaata	ccaggtgaga	accoccaago
1921 g	gtatctgcg	ccaaaatcgg	ctgaccccct	ggtttgtggt	gaatgaggat	gaaaacgaca
1981 c	gctctctta	attaacacga	taacgttaaa	taatcacacc	ttcctgccag	gtacggggga
2041 a	ggttaacta	ctctatcaag	gaaagggttt	atgactgtaa	acagaggcga	taacctgcat
2101 ca 2161 ta	aaaaaacgc	cggaagtgac	ggttctggat	aaccgggggc	tgaccgttcg	cgagctccgt
2221 a	ctctctcag	atcaattaac	acacaccacc	accgatgaac	ggatcacccg tgtttgactt	ccatcggttt
2281 g	ataatacag	tcaatcctaa	catgatttat	gatactgcac	tgaccggtga	acaycayacy
2341 a	caaggagtg	tcgatgcggg	taatgatctg	atattgaatg	acattaccgg	ccaacctata
2401 c	tggccatca	atgcaaccga	agtcactcgt	acgtggcaat	atgagaatga	cactttaccc
2461 g	gacgcccgc	tcagtatcac	agaacagcct	gctggcgaag	caggccgtat	cacagagcgt
2521 ti	ttgtctggg	cagggaacag	tcaggcggag	aagaacagca	acctggccgg	acagtgcgtg
2581 cc	gtcactatg	acaccgccgg	actgaaccag	acggacagta	ttgcgcttaa	cggcataccg
2641 ct 2701 ga	cgtccgtca	cgcgccagct	gctgccggat	ggtacggacg	cagactggca	gggaaacaat
	ccaccagect (	ggaacgaccg	getggeaeeg	gaaaacttca	ccaccctgag	cacggcggat
	acatageag	geetgetgae	taccactyat	ctacaactta	acctgcagcg cgggcgggac	rgrggcgtat
2881 at	tcgtgaaat	ccctgacgta	ttccaccaca	ggtcagaaac	tgcgcgaaga	gcacggcc
	gcgtggtga	ccacctacac	ctacgageco	gagacccagc	geettgttgg	cataaaaacc
3001 aa	aacgcccac a	agggacatgc	acaggggacg	aaggtgttgc	aggacctgcg	ctatgagtac
3061 ga	acccggtgg (	ggaacgtggt	gaaagtgacg	aacgatgcgg	aggttacccg	cttctggcgc
3121 aa	accaaaaag 1	tggtgccgga	gaacacctat	gtctatgaca	gcctgtatca	gctggtcagt
3181 gc	ccaccgggc (	gcgaaatggc	caatatcgtt	caacaaaqca	cactattacc	cactccttcc
3241 ct 3301 as	cattgata (	gcagtaccta	cagcaactat	tcccgcacct	acaattatga	ccgtggggac
	eccigacge a	ayaracgtca	cagegeeeeg	gccactggta	acagttacac	cacggacatc
3421 gt	togatocae i	ttttcactcc	aaacaaaaaa	cagatagas	tgacggatga tgcaaccggg	tccggcaaag
3481 gt	retagaege (	cacacaatas	actactass	ataaceccac	tgcaaccggg	cagaacctc
	cgaccagg a	aatcctatco	ttatgatge	accaatcaac	gcatcatcaa	aacccacatt
3601 ca	gcagacgg (	ctaacagete	gcaggcgcag	agcacactat	acctgccagg	actagaacaa
3661 ca	caccacaa 1	taaatggcac	gacqqtqaaa	gaggtgctac	acgttatcac	gataggcgag
3721 gc	egggccgtg d	cgcaggtgcg	ggtactgcac	toggagaaco	gaaagccggg	toccatcagt
3781 aa	icaaccaga t	tgcgctacag	ctatgataac	cttatcggca	gcagcggtct	ggaggtggac
3841 gg	jtgacggac a	aattatcag.	tatqqaaqaa	tactacccot	acqqqqqcac	tacaatataa
3901 ac	ggcgagga g	gtcagacaga	ggctgattac	aagactgtgc	gttactcagg	caaggagcgg

Fig. 2

chrim5ed2.seq gatgcaacgg ggctgtatta ttacggctac cggtattacc agccgtgggc ggggagctgg ctgagtgcgg acccggcggg cactatcgac gggctgaacc tgtaccgcat ggtcaggaat aacccggcga cactggatga taaaaacgga ctagcgcccg gaaatagata tgtatttttt ccatttattc atgaggacag gatttttcgt ctggcaagcg cgaatgttta cagaacggaa cataataaat ctgacatcat tgcggttgta gaagataaag cattagatag taaactattc accaatagta ttgagcagtt tttcaaaaaa cctaaaggaa aagcaatcct gaaaggatcc cctgatatta aagaaagget actcaataat atagtacatg acctgagcaa tatgcaggta ggagatcage tgtatgtaaa cgctcatggt cattctgcga aaccattttt ttactccgat tcgggatatt caaaaatcat catggaacag ctccaaagag gggctaacta tgtagctaaa gatttagtaa ataagtttaa attaccagaa aatgcaacaa tcaagataag tacgtgtcat agtgctgaag gtaagggcgc tcatattacc gtcacatcca ctggaacaaa tgaaaaaatg agatacagtt ccattataga gaacaaaggg gaattttccc ggtctttagc aggtaccatg gaaaatgagt taattaaact acagccgggc agagttcgcg ggaatgtata tggttatctt ggcgcgacaa cgttctatgg tgctaaaaat gaaaaagtca tacacctcaa agatggcaat ctgacaactg gcgttcatga aggcaagtta tcaatgttta ctaaaaagaa ccgattttca gaaaacattt ttgggttaaa ggtaaaaaga agtotgacgo gaacaaactt tacoggoago ggcgtataaa aacaaattto aaaccgoatt taaatacgga cagccagogo gtgctcaaaa cgacotgaco tgtcacgtog ttgcttccco gttacaccgg caggtoggac gcctgctggt citttacccg tittcactgg atattgaccg gcataccatg agcatcacga ccaaatgigg ttctgtcgca ttaacggcga gttaatcaga gatcccgcga gtggttgttt tttatactgc caatcgataa aattgttccg ccgccgataa acattcacct tcctcttgcg aatacgggtc tttccggage atggcgacga gttctatccc tgccagtcac gtttgtgccc ggcgaaacga tttaaaacct aacattggtc gtatccgacg tttgacatgg cgatggtctt gctcaatgcg gttattcagg tatttattt gccggatttc aatcccctgc tgaggctcac cttccgcatt gaggagggtg agcycggcgg tattggccc acttttatcg atagtcacca cagtcggtct gcccgtcggt atcaaccgac cggtaaaggt atttcaactg gcctttgact ttaatgtatg tttcatccat tcgccatcgg gagccaacag gctttttgcg tttccggaaa gccttgctga gcaggggtac caaacgtaac acccagcgag gtatcgtggc gtggtcgacg aagatccccc gtttigccat catttoctcc agattacgta agetcagege gtaggtcaga gaccaacgga cacattggge catgatatcg acaggataat ggaggagttt gaatgcgttt cggatcaggg acatcactct ggactcactc aaaaacagga gagcttacct gatattacgc taatgcgaca gaacctattt tatcgattga ctgcataaat agtcatcatt cccacctccg tacaaacttt ctctgttaat gcgacagaac cctttttcc atattgcctt gcattgcctg tagttgcctt crettetaat gegacagaac cettettee atattgeett geattgeetg tagttgeett aaatteettt aaatteett tagttgetta aaatateeat aatattgeet gageegattg gegaagegeg teatttgage eggteggega ggegeteag caaateeate egetegtgaa aaategeeac aattaaegea ggegeattt caegeggeag acaaaacaa tagtgaegt cacaattgege cattgegeaat geeggaaga gegegeteat gteettaaaa gageettgge eggacageaag cegeggaatg eettgeegga teggeggtgat teggeggtgat teggeggtgat aateggeetea geeggeteet gagatacata geeggteget eeeggaagt tetteatega gaatttegee gataetttte eegcaceet teeeeteeat eeetteattg atgeggata gatactttte cogcaccet tecetecat coefficatig atgogtgtat ccagcaagge tttcagetce tgccatgcet gategetate getcatteeg ggaaacagge gttcgagtge gtattgettg ategtettge ettgcaagge agccaatgce ttcaggettt ggtgetgeeg gtcggtgatg tcaatggtca gacggctcat ggttttctcc ttatcggtta tacacaaacc cacattagca catataatgg tttgtggcta tttattacac agggggtcag atatcgtttt gtccggcaaa cattcgtttg tcggtcattg tcattttcag aagacgactg caccaattaa tagggattag agcctggtgt gcatatgact ggcactgttg caaagtttga gtcactgatt atgctaagtt aaatgtttcg taacgggagc tcaccgtatg aatgtattca aaggccgtca ttttacaggc attatcattt tgtgggcagt tcgttggtac tgcaaatacg gcatcagcta tcgtgaactc caggaaatgc tcactgagcg tggcgtcaat gttgaccata cgactcttta tggttctgtt gcaataagta aagcctcggt aatatctcac ttatttgttg ataagagtgt cgtcaatgtc gttgatacga aaagccttca tgacacccat gcataagaga agcccatcat taccaagggc ggtattcctg ccgcctgagg gtttctgctg attatccctt tctgcctttt cccgcattct gttagaatgc ccacttctta attgtttcca aaactaatgt tgttatgtca aatcaagatc ctcataataa acgggacagt ctgttctctg ccccaatcgc caatttaggc gactggagtt tcgacgaacg tgttgccgaa gtctttcctg atatggtgaa acgttccata cccggttatt ccaatatcat ctccatgata ggtatgctgg ccagtcgctt cgtgacgcca ggtagccaaa tttatgatct cggttgctcc cttggggcgg caactctgtc catccgccgc agtatcaatg ctgataattg ccggattatc gctatcgata attcaccagc catgatcgaa cgctgccgcc gccatattga ttctttcaag gccagtacac ccgttgaggt gatcgaacag aatatccttg ataccgacat tcaaaatgcc tcgatggtag ttttgaattt cacattacaa ttcctgcacc ctgatgatcg ccagaaaata ttgaagaaaa tttacgcagg attaaaaccc ggaggggttt tggttctgtc tgaaaaattc aattttgaag accagaaaat tggcgagtta ctattcaata tgcaccatga tttcaagcga gccaatggtt atagtgagct ggaagtcagc caaaagcgca gtatgctgga aaatgtcatg cggacagatt ctgttgacac ccataagtca cgccttaaag aagtcggttt ccagcatgta gaagtctggt tccagtgttt caatttcggt tcattattgg caataaaagg aactgaacaa tgatcgattt cggtaatttt tatcaattga tegecaagea cecactaaac cattggetgg atageetgee ageacaattg agecaetgge aaaaaacate acageaeggt cagtteaget catgggtaaa aattetggaa aatttgeetg agatcaagcc aagccacctt gacctgaaaa atggtgtcat tgcgattcat gagccggatc tgtcaaaagg tgaaaaagct cgcctccaca atatcctgaa aatattgatg ccatggcgaa aaggcccttt ttcattgtat gacgttgaaa ttgataccga atggcgctct gactggaaat 

Fig. 2(i)

chrim5ed2.seq gggagcgagt gctgccccat atttctcctt tagaaggaaa aaccgtactt gatgtcggct 8281 giggcagigg tratcacatg tggcgcatgg tiggcgaagg cgctcaattg gitgigggta tcgatccaac ccaactttt cicigtcaat tigaagcgat cagaaagtig tiggggaaca 8341 8401 atcaacgage ceaecttetg ceattgggea tegaacaatt accegaactg caageetttg 8461 8521 atacggtatt ttcaatggga gtgctctacc accgccgctc acctcttgat catctgtggc 8581 aactgaaaaa tcaactggtg tctgatggtg agttagtgct ggaaagttta gtgattgagg gtgatgaaaa tcagtgcctc attccgggtg aacgctatgc acaaatgcgg aatgtctact 8641 ttattecete ggecaagatg etgaaagtet ggetggaaaa atgtggtttt gtegatgtea 8701 gaattgtcga tcatgcggct acaacacctg atgaacagcg ccggacagaa tggatgaaga 8761 ccgaatcact ggtagatttc cttgacccat cagatcacag taaaacaatt gaaggctacc 8821 8881 ctgccccatt gcgtgctgtc ctcattgccc gcaaaccata atattgaata aatattaatg agtgaactgt ccaatatggc aattcactca ttaagttcta agatttcgct ttccttatga 8941 cgcaagcgat atcacatcta ccgcttaatc aggctcatca ctcccttcat cgactcaact 9001 9061 aactcaccat caacactgta gtgagaatat tcatctacat cacgatcgtc cgaactcatc gccaccccta cctttcggta cttcatggta gaaggtacaa ttttacccgc cgagctatga 9121 acttccgtca ttccggcttc cagaaactta ctgatattac tcaccctgac acccgcgccg 9181 9241 ggcataatta tcggcccacg gctggcttgc atcagctctt ttaacagcgt cagccccagt tcagcattct gctgttggcc tgatgttaaa atacgctgca ctccaagctc tgtcagttgt 9301 9361 tccaacgcaa catgcggatt aaaacacata tcaaaagcgc gatgaaaagt aacagccata tttcccgaca gtgacatcaa ctgccgcata cggagtctgt caatatggcc gttttcgctc 9421 9481 9541 9601 9661 9721 tacgaattcg tcaacaagtc atcatttaac ctaaaactac tttattagtg aacttattaa ctatgacaac taacttatct taatgacatg ttggaaatca caaggtcaga atttttact ggaacagcat gataaaaacg tcattttgc cggctatacc tcacttttga caatttctct 9781 9841 9901 gatacaaaaa ggatgatatt tgatcgtgat ttcaccatca gtgacagcca aaatcgggtt 9961 10021 cggcaaccgt tcattttttc cttttggatg actctccttc atccgaaaat tatctggtaa aggttgaggc aataactttt tagcttcaca aataagttgc tctggtggta caggcagcac 10081 cagttcgaca tgttcccaac cttggtggac atagtgtttt tggctgggat aaggcaattc 10141 cacgcaatca attttccagt caagtagtac tattggctgg ttcagatcaa ataagcagat 10201 cggacggcca ttaatgatac tttcagatat taactgacca cattgcagaa agcccttacg 10261 ccaacgatcg gctattttac tttcattaca acgcagggaa atatggtctg ctgaatattg 10321 10381 ttcgaagtgt aagccaagat gcccttcaaa ttgcttaagt ttttgctcaa atccggctaa atcagccact aaatcctgta actcagaaat ttttgaaaaa tgagacattt cttccccatt 10441 10501 aaataaaccg taaaaattgc gttccattaa tgtgacataa atcacaaatc gtctattttt gccgaaatat cactcagtaa gcgactaatt gaagttggca taacgacgaa tcgcctgaaa 10561 10621 gacaggetaa aaacaaaaag taacaccace agaggtgget gatggtagca tgcaggacce cgaatatggt ataaaccccg ttattttete ataacccaca cgettaaaag tattgcattt 10681 ccaaaatgca taagctttcg tgcgtaactt aaggtaacac ggtgaatata caggttattc tttcagaaaa aatcagcaat gcgctgattg aagctggcgc tccaaacgac agtgaagctc acgtccgtca atctgccaaa gcacaatttg gtgactatca agcgaatggt gtgatggctg 10741 10801 10861 ccgctaaaaa ggtgggaata cctcctcgac aattggcaga aaaagtcgtc agccaactgg 10921 10981 atctgcaagg aattgccagc aaagttgaaa ttgcaggccc aggttttatc aatatttttc ttgataaagc gtgggttgca gcaaatatag aaactaccct gaaagatgaa aagctcggta tcaccccagt ggaaccgcaa accatcgtta tcgattattc cgcaccgaat gtcgccaagc 11041 11101 agatgcatgt tggacacctg cgctcaacca tcattggcga tgctgcggcg cgtacccttg 11161 agtttcttgg gcataaagtt attcgagcca accacgttgg tgattgggga acccagttcg ggatgctgat cgcctatctg gaaaagatcc agaacgaaaa tgccaatgac atggcattag cggatttaga agctttctat cgcgaagcaa agaaacacta cgatgaagat gaagagtttg 11221 11281 11341 ctattcgcgc tcgtaactac gtcgtcaaac tgcaaggcgg tgatgaatat tgccgtaaga tgtggcgtaa gctggtagat atcaccatgt cccagaatca ggaaacttat aaccgcctga 11401 11461 atgtcacatt gacagaaaaa gacgttatgg gtgaaagcct gtataacgat atgctaccgg gtatcgttgc agatttaaaa caacgtggaa ttgccgttaa gagtgatggc gcgacagtgg 11521 11581 tttaccttga tgaattcaag aataaagaag gcgaacccat gggcgttatt atccagaaaa aagatggtgg ctatctttac accacgacgg atatcgcctg cgccaaatac cgtcatgaaa 11641 11701 11761 ccctaaatgc cagccgtgtg ctttactaca tcgattcacg ccagcaccag cacctgatgc aagettggge aattgtacgg aaaacgggtt atattccaga atccatgtca ctcgaacacc 11821 11881 acatgtttgg catgatgctg ggcaaagatg gtaaaccatt caaaacccgt gccggcggca cagtaagact gtccgatttg ctggatgaag cgattgagcg tgcggatacc ctcattcgtg 11941 agaaaaaccc agatatgcca gaagacgaac tgaaaaaagt cgtggaagcg gtagggattg gcgcggtgaa atatgcagat ctttccaaga gccgtactac agactatgtt ttcgactggg ataatatgct ggcctttgaa ggcaacacgg caccttatat gcaatacgcc tacacgcgcg 12001 12061 12121 tgtcatctat ctttaaacgt gcggatatcg atgaaaacag cctgacactg ccggtgatgc 12181 tgaatgaaga acgcgagcag gcattggcaa cccgcctgtt gcagtttgaa gaaacgatca ctaccgtcgc ccgtgaaggt acgccacatg ttatgttgc atacctgtac gatctggccg gtetgttctc tggtttctat gagcactgcc ctatcctgaa tgccgatagc gaagaactgc 12241 12301 12361 12421 gccagagccg cctgaagttg gctctgctga cagcgaaaac tttgaagcaa ggtcttgata ctctgggtat tcagactgta gaacgtatgt aataatcttc tacaaagctg aaaactggcg 12481 tgatattatt ttattacgcc agttttttcc tttcttctat tctgtcaaaa attaccaatc 12541

Fig. 2(ii)

			3/14			
chrim5ec	i2.seq					
12601	tgacatataa	ttaatcttga	gctaacattt	tgatatttaa	tcatagaaat	atatgaacac
12661	agaagcagtt	gttatgaaaa	aattatttat	ttctaactgc	tagccctacc	aatctccata
12721	aaaaaacctg	atttttattc	actattaata	attaatgata	atttctattt	taattaacct
12781	tgttataaaa	aatagtattt	taaaaaaaca	ttttacatta	tataaaatat	atcaatcgac
12841	tctttatttc	tttatccatt	tataaaatat	attttttacc	aaaataatat	ttaaatcata
12901	tattatattt	acatcacgtt	agatcaaaat	aacaattttt	tagtcgttaa	cccagattca
12961	gaacggcacg	atgatatata	agtcacatgg	gttgtaaata	aaggaaaaag	ataaaaaaga
13021	ggagataaat	cttcgcattt	cttcaatgaa	gatggatata	gatactgtaa	ggtagtaatt
13081	taaatgaaat	ccattaacaa	attaatattt	aatttacatt	aaqagaggat	tctatgagtg
.13141	attonacton	totttcaaca	tttaatotta	ttcttgaaac	aggattagat	aactgcaata
13201	totacoctaa	toggcttaac	atgattgggg	taattattaa	tatcacaccc	actgatgatg
13261	aagggaactt	cotagatatt	gacgatgtta	cactaaatga	taacatcaag	attgttgatt
13321	atatedated	aagcgacatt	gatogcagto	acquatqutt	ttatacagga	aatcctaatg
13381	aatacaacac	tattccaaat	agtcagtctt	attctttatt	aaaqagtgaa	aattctcaaa
13441	ttacgcaaat	taaacgatat	atttcttatt	caaatacatc	caggctaaga	accaagtctt
13501	tttctgcgaa	ggtaaccact	accagtggaa	aagttatttc	aataactcaa	aatagcatta
13561	attcatctcd	ggtagtaatt	aatocaatao	atgcaactaa	ttttactgat	gatgaacttc
13621	Caacaacaaa	agaaacaagg	tttgaaaatc	aatcctatac	gtcacataaa	tcatctacaa
13681	actetttata	tatacataca	togacaatac	caagaagctt	aaaactacaa	aattggcgtt
13741	gggaagatta	caataatggc	tagacttaga	cacaaagttg	ctactataaa	acaggagccg
13801	atogaggato	agagtcaacc	cgctggttgg	ccgctggttc	aatctttcca	ccaggaaatt
13861	atmatmmcct	gtggctagat	aatgatatcg	cactaagtgg	tatggcacac	aaaagctaca
13921	atottoatac	tootatcaat	caattgagtt	ttacccqtat	tataggtaaa	ggtttcagct
13981	aagtttataa	tatatccgga	cttgatagag	agcataccat	tattattatc	gaccagtatg
14041	otaacaaata	tagaatatta	ttccatgccg	ggtatgaaaa	ctcagatcci	tacttgtctt
14101	catcaatagt	atattaaaaa	tagtgttacc	aataatataa	tagcatttcc	ccaatacgaa
14161	acttmaccta	gtggctgtaa	ttataattta	ttttcacage	cactttttaa	gtatacataa
14221	acttgattea	gttattcagg	agaatcacca	tgagtgacaa	taatgagttt	tttactcaag
14281	ctaataattt	caccageget	gtcagtggtg	acattaacce	tcgcacagga	ttatacaata
14341	tacaaattac	tttaggtgac	attgttggta	accotaatct	tggacctact	ctgcctctta
14401	ccttaaccta	ttctcctctt	aacaaaacag	atattogatt	tggcattggt	tttaattttg
14461	gattatcagt	ctacgacaga	aaaaattctt	tattgtctct	ttctaccggt	gaaaattata
14521	aartcatcga	aaccgataaa	acagtaaaac	ttcagcaaaa	aaaactcgac	aatttacgct
14581	ttgaaaaaga	cctasaagaa	aattottato	gtattataca	taaatccggt	gatattgaag
14641	tattaactaa	tttcaataac	aatgcctttg	acctgaaagt	ccctaaaaaa	CTATTAAACC
14701	ctactagaca	toctatctat	attoattooa	attttgaggc	aactcaacct	aggetaaate
14761	gtatttatga	toatctocat	gggcatgata	taccattatt	aaacctagaa	tatcaaggac
14821	teattasaac	gatattaacg	cttttcccta	ggcaaaagga	aggctaccgt	accgagctac
14881	getttetaaa	cagacaattg	aacagcatcc	acaactttag	cttgggtaat	gaaaaccctc
14941	tracttootc	cttcggttat	acccctatao	gaaaaatgg	tattttgggg	caatggataa
15001	caagtatgac	cactcctaga	ggattaaaag	aaacggttaa	ttatagtaat	aataatcagg
15061	ggcatcattt	ccccaatca	occaatctac	caatattacc	ctatgtcaca	ttaatgaagc
15121	aggttcctgg	agcaggacaa	cccactatac	aagcagaata	ttcgtatacc	tctcataatt
15181	atotogotog	gggatctaat	ggtatatgga	ataataaatt	agataatctg	tatggattga
15241	tracagaata	taattatggc	tctactgaat	cccacagata	taaagataaa	gaaggccatg
15301	atcaaatagt	ccatatagaa	cocacataca	ataattacca	tctgttaact	tccgaatgta
15361	agcaacaaaa	togatatata	cagacaectg	agacagcata	ttatgctatt	attggccata
15421	attttgattc	tcagccctca	caattccagt	tgccaaaaac	caaaacagaa	acttggcgta
15481	ntacagataa	cagctatega	agtgaaatta	ctgaaaccac	atttgatgaa	agcggaaacc
15541	. ccctaaccaa	agtaatcaaa	gataagaaaa	cacaaaaaat	aatctcccc	tcaacgcatt
15601	gggaatacta	cectecaact	aaaaaaatca	ataattoccc	accagaaccg	tatggattta
15661	ctcatttcat	aaaaaaaatc	atacaaactc	cctatgactc	cgaatttaaa	gatgatccgg
15721	aggaatttat	ccagtatcgt	tatagcctca	ttaacaatca	qaqtcatgtg	actttaaaaa
15781	tagaagagcg	ccactacagt	gcaactcaac	ttctgaatag	tactctattt	caatataata
15841	cogatasaso	toaacttoot	cotttattaa	aacaaactga	atgtaccaaa	ggagaaaatg
15901	gaaaaactta	ttctatcata	cataaattta	cctatacaaa	acaggacgac	acgctgcaac
15961	agagccattc	cataaccacc	catgataatt	tcacaattca	ccgcagtcag	gttcgttccc
16021	gttataccgg	acatctattt	tctgacacag	atactaaaga	cattgtaact	caaatgtcct
16081	atoacaaatt	aggtcgatta	ctcacacaca	cccttaattc	cggtacacca	. tatgccaaca
16141	ctctgacata	tgattatgaa	ctaaataatc	ttcaggatga	caatcgccct	ccgtttgtta
16201	ttaccaccac	ggatgtaaat	ggcaatcagc	ttcgcaatga	attcgacggt	gccggacggc
16261	atotcaocca	atocctoaaa	gactccgatg	gtgatggaaa	attctatacq	atacatacgc
16321	ascastatos	tgaacaaggg	catcatcata	catctacata	ctccgactat	ctcacaaatg
16381	Tabas cases	gacggatect	gataaggtgc	atctotctat	gtcaaaatco	tatgataatt
16441	gaagacaaca	tacasacaca	cactogagt	atggggtttc	agaaaaaata	actgtagatc
16501	castascatt	dacddcaca	Asscartter	anagcantag	caataatoto	caaacgggta
16561	agazaatatt	aacttatace	ccaartcaac	aacctataca	gattacgtta	tttgacgaag
16621	aayayyttat	acadeatect	cataccetes	ctcacata	ctagnatann	gttcgcaaag
	nange cast co	acayayııyı	tacactett	accastates	taactataac	cgagtcattc
16681	adaccgatge	toctorter	accatests.	atcaatatya	+00200000	agtactgata
16741	aaacaacgct	nantation-	accategtaa	tttoctto-	ecarcaeaco	tttgacgggt
16801	cgctgataac	agatattega	grydarggaa	ancanatata	acaycaaacy	tattcggcag
16861	cgagccgact	aacacaaagt			gycciacaci	. June Coggody
			Fig 21	ni)		

ch rim5	od2 coa		0/14			
	ed2.seq					
16921	gtaatgacca	a atgcccatca	acagtaataa	caccagatgg	tcagtttatc	cattatcaat
16981	atcagccaga	a attagatgat	gcagtattac	: aagtagcatc	aaatgaaatt	actcagcagt
17041	tcagctataa	a cccagtcact	: ggggcattat	taaaggcggt	ggcagaggga	caaagcttga
17101	cccctatcta	a ttatccatcg	ggaagactta	agatggaaaa	tatcaatgat	atgaaaaaaa
17161	tgagttacct	: atggacactt	aggggtctgg	agaacggtta	cactgatctg	actogaacaa
17221	tacagaaaat	ttcgcgtgat	acceatooca	gggtgacaca	aattaaagat	tcatcaataa
17281	agactactct	asattacoat	acctastc	accetettaa	tagtcaagta	zezezeteze
17341	caactaate	tatattaaca	. gaaaaagaaaa	32tttastaa	cttaaaccga	acayacccay
17401	agactygete	taryrryaca tartrastar	acaacaytyy	tacttgatgg	cccaaaccga	gaaattggac
17461	ggaaactgt	, cyatayetta	ggccatacgt	cagacaceca	gcagagctgg	ctgaaaacac
	aycaactayc	aaacagaaca	gtgaaactga	acggagtatt	gcagcgtaca	gaacagtact
17521	Cttacgattc	ccgtaatagg:	ttgaaccaat	ataaatgtga	cggtgcggaa	tgcccgacag
17581	acaaatatgg	, ccatagcata	gtcacacaaa	attttactta	tgatatctat	ggcaatatca
17641	ccgcctgtca	caccacatte	gcagatggga	cagaagacca	tgctaccttc	aaatttgcca
17701	acccaactga	cccatgccaa	ctgacagagg	tacaccacac	tcatccagat	atgccggata
17761	atatcaggct	: gaaatatgat	aaggctggta	gagtaataaa	tatcactgat	aaccatggaa
17821	atacggaaaa	ctttacctac	gatacattgg	gcagattaca	aaacggtcaa	ggtagtgttt
17881	atggttatga	tccattaaat	cocttaotoa	gtcagaaaac	agatacccta	gattgtgage
17941	totactatco	ggaaaccatg	ttggtcaatg	aagtacgcaa	tggagaaatg	atcoatttat
18001	tacqqacqqq	toaaacaata	atcocacao	aacococatc	aaaagtcttg	ctanceccae
18061	cagatageca	acadadeeta	atetteecas	atastasaca	aaacttgtct	ccaacayyaa
18121	atantocata	taassaast	acatcaacga	gryatasaca	ttctatcctg	caagaagcac
18181	atagegeaca	teresent	adacciacay	caaacgacgc	tectateetg	ggctataatg
18241	atocases+	- cyacccagtt	aguggagtaa	cacatttagg	taatggttac	cgctcctatg
	atecaacatt	aatgcgcttc	catactccag	atagettaag	cccctttggt	gctggaggga
18301	ctaatcccta	ttcctattgc	ttaggagacc	caattaatcg	ctcagaccct	tctggtcatt
18361	tgagttggca	agcatggaca	gggattggca	tggggatcgc	tggattactg	ctgaccatag
18421	cgacaggtgg	aatggcaatt	gcagcagcgg	gaggtattgc	ggcggcaatt	gcttccacct
18481	ccacaactgc	actggcattt	ggggcactga	gtgttacatc	ggatataacg	tctattctta
18541	gcggtgcact	ggaagatgct	tcaccgaagg	catcttctat	actcggatgg	gtttcaatgg
18601	gaatgggtgc	tgccgggtta	gctgaatcgg	ccattaaagg	tggcaccaaa	cttgcgacac
18661	atctaggagc	attcgctgag	gacqqqqaaa	acqccttact	taaatcgact	tccgaaagtt
18721	ctagaataaa	gtggggagtg	acaagaagct	tagatagaga	aattgttcgc	aatgaagaag
18781	gtcaggtgat	aaaagatcat	agccgaggtt	ataccoataa	ctttatgggg	aaaggagagc
18841	aggetatatt	agttcatgga	gataaagatg	gatttttgta	tcatacagaa	desaces c
18901	ataatggaaa	agggccatac	actcgacata	ctcctgaaca	actcgttgat	tatttmaaan
18961	acaataacat	cottoatctt	acacaaggag	Gagacaaacc	tgttcattta	ttatcctact
19021	atggaaaaag	cancontaca	acadataaaa	tagazzate	tatcaacagg	ccaccccgcc
19081	cttattctaa	taaaccaaca	atatorona	cygtaaaata	cattaga	ccagccaccy
19141	tottacccaa	taaaccaaca	acaccacaag	garragecag	aatagaaaga	aaggactttt
19201	ccccaaaaay	cacciaccac	tegtatgate	cacggaagat	catactggga	agaacagaaa
19261	aaacaytyaa	accaaaaact	tttcgccct	aataaccttg	caaaattcaa	aggtactggc
	agaagtcata	taaataactc	tatgactggg	taaatgaaat	cagtattcaa	acattaaaca
19321	ggatggaagt	ggtcatctta	tcaaccgccc	aataaaaaat	tgggcggttg	tattgaataa
19381	aattatttat	taatcaggga	atcactgcaa	tccccgatga	gcaaaatctt	tcagtctaaa
19441	tcccatcaag	gccagaactg	caaaataact	gcctgctccc	gcaatcacta	cgcccattaa
19501	gcgcagtagg	cgcattgcca	tattgccctg	ttcccatgct	ggcataaccc	acaatactgc
19561	caacagcacc	ccgaccatca	cagcaattgc	caccaccaat	ttaaacagga	atatccccca
19621	tcccgccaaa	ggcgtgaaaa	tatcacgctt	acgcagttgc	caataaagca	tactggcatt
19681	gaagcaggca	gccagaccaa	tagaaagcgc	cagacctgca	tgtttcaacq	ggccaacgaa
19741	agcaaggttc	atcaattggg	tcagaatcaa	ggtcgcgatc	gcaatttta	ctagtatttt
19801	gatatettga	cotoaataaa	agcccggagc	gagaacttta	accacaatca	accccatcaa
19861	qccaaaacaq	taggcaatta	acacccacta	agtcatctca	gcatcaaaag	canaaaanto
19921	accatattoa	aataatgata	ccatcagaga	ctcccccaga	ataccgagag	caactgcag
19981	aggcaacgcc	апсавпавас	agagacgtag	ccccaatcc	atcagttttc	catctycaca
20041	gtgattacca	ctooasasa	ttttccccec	tassages	aaaatcgtcc	etanesses-
20101	acceatece	ccagaaaaac	attocattas	cyaayycayc	taatacatcc	CLaacgecae
20161	acctasseca	acasatese-	cananttet	attantant-	aaggaaatct	acgaaacaga
20221	tagagaga	ayaaacyagg	caaaaaccyc	accaacgace	aaggaaatet	gcccgaccga
20281	cacacacaga	arrigeaggee	ccatttgacg	gataacccgc	catacggcac	tgtcacggaa
20201	agaaatccgc	ggcaatacca	gcatgccgat	ctttttcaga	tgaggaagct	gataggccaa
	ttgcaaaacc	cctccggcaa	caacggccca	acccagcgcc	agcactggcg	gattgcaata
20401	aggagccaca	aacaatgcaa	aaatgatcat	actgacattg	agcagtgtcg	gagcaaaagc
20461	aggcaccgaa	aagcggttcc	atgtattaag	aattgcgcca	gccaaagaag	ccagcgaaat
20521	caaaaagata	taaggaaacg	taattctaag	taaatcacgg	gttaagacaa	acttatccoo
20581	tgtatccgta	aatcccggcg	cagtcacata	gatgatccaa	ggtgcagcaa	ttacacctat
20641	cactgagacg	atagccagaa	tcaatgtcaa	catacctgag	atatatacaa	taaaggtacg
20701	tgttgcttca	tccccttgtt	gatttttgta	ttcggcaaga	ataggaacaa	aagettgega
20761		****	tacoocotaa	caggttaggt	aatttaaann	Caacaaaaa
	aaaagcgccc	LULUCAAAGA			aaayy	Juacaaaaa
20821	aaaagcgccc	gccattccta	caccasatat	acmamcaat-	2+444c2+44	// D D D D D D D D D D D D D D D D D D
20821 20881	aaaagcgccc ggcatccgtc	gccattcctg	caccaaatat	acgagcaata	atggcatcac	gaataaagcc
20881	aaaagcgccc ggcatccgtc cagcacgcga	gccattcctg gaaaacatcg	caccaaatat	gaccgctgcc	agtgatttca	ataagttcat
20881 20941	aaaagcgccc ggcatccgtc cagcacgcga ggtattgttc	gccattcctg gaaaacatcg taaagttgta	caccaaatat tcattgaact ttcttatgga	gaccgctgcc attaagcata	agtgatttca aaaatgtaaa	ataagttcat
20881 20941 21001	aaaagcgccc ggcatccgtc cagcacgcga ggtattgttc caggcatcat	gccattcctg gaaaacatcg taaagttgta aaaaatggca	caccaaatat tcattgaact ttcttatgga tataaagcaa	gaccgctgcc attaagcata tctggcggga	agtgatttca aaaatgtaaa tagcagccgg	ataagttcat gctatcccat tgtttaaagt
20881 20941 21001 21061	aaaagcgccc ggcatccgtc cagcacgcga ggtattgttc caggcatcat ctaacagaca	gccattcctg gaaaacatcg taaagttgta aaaaatggca aaaaccctgt	caccaaatat tcattgaact ttcttatgga tataaagcaa ctacattttc	gaccgctgcc attaagcata tctggcggga tatattacgc	agtgatttca aaaatgtaaa tagcagccgg cccattagcc	ataagttcat gctatcccat tgtttaaagt ttaccccgag
20881 20941 21001 21061 21121	aaaagcgccc ggcatccgtc cagcacgcga ggtattgttc caggcatcat ctaacagaca attcataaac	gccattcctg gaaaacatcg taaagttgta aaaaatggca aaaaccctgt ccactttgcc	caccaaatat tcattgaact ttcttatgga tataaagcaa ctacattttc ccatgcatcg	gaccgctgcc attaagcata tctggcggga tatattacgc tcagtcaatg	agtgatttca aaaatgtaaa tagcagccgg cccattagcc	ataagttcat gctatcccat tgtttaaagt ttaccccgag
20881 20941 21001 21061	aaaagcgccc ggcatccgtc cagcacgcga ggtattgttc caggcatcat ctaacagaca attcataaac	gccattcctg gaaaacatcg taaagttgta aaaaatggca aaaaccctgt ccactttgcc	caccaaatat tcattgaact ttcttatgga tataaagcaa ctacattttc ccatgcatcg	gaccgctgcc attaagcata tctggcggga tatattacgc tcagtcaatg	agtgatttca aaaatgtaaa tagcagccgg cccattagcc	ataagttcat gctatcccat tgtttaaagt ttaccccgag

Fig. 2(iv)

			1117			
chrim5ed	2.seq					
21241	ttgagcaggt	tcccactacg	atggttatgt	ggcatgtaat	tagccataat	ceggegeatt
21301	ctctgatage	gctgcatata	gcggatcgca	ggttgcagcg	ecteatacat	accacataca
21361	ccttgcaatt	tctcagctaa	agaataacca	teacacteat	cagagtccaa	taatacaaaa
21421	ttaaatgtcg	cattetacca	ttotaactca	gcgatatagt	tcaaccgatt	totccatcca
21481 21541	gagggttgat	assactocc	atccettatt	gactacage	gcttcaggaa	atctttattt
21601	tctgaataat	actcagccaa	tcgataagca	tcacottcat	ataccagacg	aacaaccatc
21661	ctatecetaa	taaatcqaat	tttgggcgat	gcagaacgat	aaccaaacat	gccttctcct
21721	gatatttgtc	ttgaaaaaat	tctggataat	ttatctctta	ttaaaattat	tgcctattac
21781	cctacgataa	aaaaatatca	tctataaccc	tctaccttaa	agatgagatt	agggtcagaa
21841	taataagaac	ttcatattta	attctctcat	attttagtgg	tgcgcaatgt	cgttggtttc
21901	acaagcacgt	agcttgggta	aatacttcct	gctatttgat	aatttattag	tegetetagg
21961	ttttttttc	gtttttcccc	taatttcaat	tegtttegte	gaacaacttg	tagagatett
22021	attgattgtt	ggtttcgctc	rggggcttcg	canaccasta	cagcaaggct attgttactg	gcatgttatt
22081 22141	eggeggegee	actiguagact	tratorcast	nacacataaa	ccatggatat	tactactttc
22201	ctacattcta	tcaggattgg	gaggaacatt	gtttgatccc	cccagagcgg	ctttggtcat
22261	taagttaacc	catececata	agcgagggg	tttttattca	atcctgatga	tgcaggacag
22321	cacaggtacc	ataattaaca	cactcatcgg	aagctggttg	ctgcaatatg	atttcaatat
22381	catctactaa	attootocat	ccatttttqt	gctggccgca	ttatttaacg	cctggctact
22441	acctacatac	cotatttcaa	caatccgtac	tcctatcaaa	gaaggcatga	tgcgggttat
22501	tagagatcgt	cggttccttt	actatgtgct	gacattgacg	ggttattttg	tcctgtcagt
22561	acaagtgatg	ctgatgtttc	cgattattat	ccatgaaatt	accggcactc	ccatcacaca
22621	caaatggatg	tatgccattg	aaaccgctat	eccectgaca	ttgctctatc	ttttaataaa
22681	ctggagtgaa	tttcccatcc	gactggagca	tcaattacat	gcgggcttat acactgtttg	acctacttta
22741 22801	cctcttttat	ttaggtttgg	taacagccga	tectacteat	gaaacgctga	gtgcttcact
22861	gtctgatcca	caaacacata	gcagttatat	gggatttagc	cgtctgggtt	tagctctagg
22921	taatacaata	ggttacaccg	ataaaaaata	gctctatgat	actggccgtg	acttgaatat
22981	gccgcaatta	ccctagattt	toctaggatt	atctaatttg	attaccattt	atgctcttca
23041	tegecaatte	aatcagaaga	aaattgatcc	totoatoctt	ggtagacatt	aatcctatta
23101	gaaatagtgt	aaaatctgcc	cattgaaaat	aaaaaggagc	caatatgaaa	agacttca
23161	taggggcagc	attagtgctg	graggarrag	testsaaaa	tgatcaattt agtgcattat	cagaaaaaaa
23221 23281	gcatcaacga	aggicetgatg	aatgattatt	tracactaga	ggatttatcc	agccagatag
23201	accaccaaaa	tectgaaaaa	attgaactat	ccacacaaac	aaaagttcaa	cttgcaacac
23401	tactgggtac	gattcaggct	gatatgaaac	tcactatcaa	ggctaaaccc	gtatttgatg
23461	CACAAAAAGG	coctattttc	gtgaaagggc	togaaatcgt	agactaccag	acaacaccgg
23521	aaaaagcagc	aactccaatt	aaggcattga	ttccttatct	gaatacctct	ttgagtgagt
23581	ttttcgatac	tcatccaatt	tatottctoa	atccagaaaa	aagcaaggcc	gaggcagcag
23641	cctcacaatt	cgctaaaagg	ttggaaatta	aacccgggaa	gttagttatt	tratttacco
23701	ataaataatt	tttatcgtca	regaaataat	tegestatte	tattaatgga atggccttcg	gctgaactat
23761 23821	tatesteece	ctatcactca	gattctaatt	catcacgcaa	tgcctgcaac	gcttcacgag
23881	traceators	cagcottcoa	tasasaccac	toottocato	ggcttcaact	ttaccgagga
23941	actoccaca	ccaaggcaaa	agatactcat	caaacaacgt	aatttgtgct	tggatttcat
24001	cttcagctgc	ctgatcttcc	agccatgaca	caqccagcaa	tagagagccg	aactgatcag
24061	caggagtate	agacagtggc	attccccott	ctgtcaaaaa	ctgacgtact	tcactttctt
24121	ctccgtccgt	ataatcagat	cgacaaacag	caacactcgc	cgattctcca	gtaaacagag
24181	cgtgataatc	ggctgccatc	actgacaatt	cactgttttg	ctgtaaacga	ctcagccatt
24241	catcctgttc	caaaggccat	gettgtttta	tangentat	agcaatcatg gcccaatatg	caacaaaaa
24301	gttgcaaaac	gggatettge	tgattcaatc	ccastaast	atcaaatagc	gttattgtcc
24361 24421	ctagadaaccc	accetactat	aaagcaactt	gttaatagcc	acgtacccta	tcgacgaggc
24481	canttoncat	ticaccetat	accatecoce	ggatattttc	acagatgaca	tccatcgcag
24541	cctcaggaat	agtottagca	gcaatatggg	gagtcacatt	aatacqaqqq	tgagtccaga
24601	atgagtacat	atttgacaat	aattetteaa	caaacacatc	aagggttgct	cctgcgatat
24661	aacctttatc	aatagcaacg	agcagatcct	gttcgacaag	ctgcgctcct	cgcgccagat
24721	testoscata	tgaacccgat	tttagttggc	tgaataatga	cagattaagg	ataccacgag
24781	tgtcgggagt	atcagggaga	aggttaatca	gcaccttgca	gccagaaaga	aaatcaccca
24841	attgctcttt	accgtaaaaa	ctttcgacac	tategagttg	ttttgacgtc	ergetteaac
24901	aacgcacatt	aaaatcaaat	tecattaatt	cettateed	acttcgccct agcaataggg	ttccaaagg
24961	CCCCTABAAC	geegacaaca	tattcatcca	tacaacaaa	ataatgcagc	actgaggtaa
25021 25081	tagestatte	ctocatctoc	cttcccattc	ctotatette	gaggcggatc	aatggaacat
25141	cagccagtaa	cataccaga	tttttcgatt	cttgtttgaa	aatcgcatcg	acaccggcac
25201	ccagtgcgaa	tatgcctttt	agcccctgac	gattggctaa	. catttcataa	ggtggctgcc
25261	aaactaatoc	atagtctgcg	gactcttgat	caccactgac	: ccattgcctc	actttggcat
25321	caggcagacg	tocctoaatt	ccttgaatcc	actcatcggt	. attaaaggaa	gggtgaaaaa
25381	aaataatott	catacaatct	tattccttat	tatttttta	atagggttaa	cctattgcta
25441	aaatoottoc	aagcctctgc	tggaaagcga	ggaatgaaaa	tattaataat	gttacccagt
25501	taacatattt	accactgcat	attacaaaaa	gcgcatcggc	: tttagtaaat	gactatcgaa
			E:- 0/	'\		

Fig. 2(v)

chrim5e			0/14			
25561	tattcaaatt	gttttttatt	tgtgtaatca	gtcaaaaagc	ctgaaaaaat	cgtcataagc
25621	ctgttgacgc	ctgccctgct	tttccctata	gtagcgcccc	gttgcagcga	cgaactcaag
25681	tgatatcgct	acaacaacaa atacgtgaaa	aatacggtga	ggtgtccgag	aggetgaagg	caccatatto
25741 25801	gaaagtgtgt taagaaag	cctgaacaca	atactaagg	ttttttataa	ttactcttga	tagaggattg
25861	aatctateat	ttagtaaccc	ttacaaaaa	tccctgacag	gacgaccgca	gaaacaaaac
25921	accordant	gtccgagagg	ctgaaggagc	acacctagaa	agtgtgtata	cotoaaaaco
25981	tatcgagggt	tcgaacccct	ctctcaccac	catctttcaa	gagaaageet	gaacttatgt
26041	tcaggctttt	togcatttat	actccccaaa	gtataggtga	aaaacctcgc	aagggttcac
26101	ctcaacaacc	tgctctaatg	ggaatgtctt	aaagatttgt	ggcttaatta	caccttgttc
26161	tagcctaatc	caaaggacat	acccgaataa	tatgaccatc	gggatcttca	gcaaggaaag
26221	tacggccaaa	aacttcggta	tggggttcct	gtactatttt	gatctcaggg	ttctttcgcc
26281	attetteaaa	gcaccgátca	acatctttac	cggatggcag	cataatacca	atttcagaaa
26341	atcgcggaat	agcacgatca	ggctttgctc	ctccactcca	aatagcaaac	agtacttcac
26401	caccagcagg	aaatgccaca	taacgggagc	tggcaaatat	cggttcagca	ttgaaaattg
26461	ttttataaaa	agcggttgaa	ctctcgatat	cagagacgta	aacaagctga	agattgggtt
26521	tgggagatac	tgtctcaacg aaagaaactt	acagatttca	gccttaatac	atttacatca	agatettgeg
26581	taccttgctt	ctgaacacga	coggectgaa	tattttcatc	totastastt	tastatcas
26641 26701	atattaatat	tcccatttga	ggaaggttaa	cctgatcggt	gaattettta	ttcacttcga
26761	tatttgaaag	ataaaaacga	atttccccca	ttccacttaa	tatcattcto	ccgtattgac
26821	ccgttttgac	ctccccttca	aattgaaaat	attcaagatc	gacttcccat	tttttccgca
26881	aatcaaaatc	gacatagtgc	gcccagatat	gggatggttt	agcgtttacc	gatgtagaaa
26941	aattgagtgt	taacatgaca	aaactctcac	agttaaatgt	atcacattga	agattaagcg
27001	agctttttaa	tacagtagaa	cagcccgcag	agtatcaatg	gtgagtgaca	atttctgtca
27061	gtagtcttta	tttggctaac	gagaaatttt	ttccattgct	ctccattctt	tgagcaatac
27121	ttgccttgaa	cgggggtaac	attggttttc	aattttcaaa	cgcatgattc	tatctgatct
27181	gaaatgacga	aattcctcgc gcaaatggcc	gtaattcaca	ttetestese	acaatgctga	tatcccaaa
27241 27301	acageetaag	attttatgcc	acactettast	cacctaacat	atctactaaa	tetecadata
27361	agcgagggcg	gctgtcgcag	cannecenat	cagtaaggag	cttocctcca	acatttattt
27421	caattctoct	gggatcacag	ccccaatttt	gcttattgca	ttatttgcag	attetttag
27481	ttagagatet	gcacgtttag	ccacccaatt	cgcgcccaat	gccaacgcct	ctatttcatt
27541	ttqtqtaaac	atgagcggtg	gtaacacaaa	tccaggcctc	aaaacgtatc	ctattcccgg
27601	ctcaccttco	ataatcgcgc	cttgagcctg	caacqatqca	atatcccgat	acagtgttct
27661	taagctgata	ttcaatttct	gcgccaacac	ttttccctga	accggaaagt	gataacggcg
27721	caatatttcc	atgagaaata	acaaacgctg	tgctctagac	aattcgtccc	atccattcaa
27781	gttaaccgac	tcatcaacct	ctaataatac	gcaactatca	taatttaatt	gattaaaaag
27841	atagttttt	gatcccttgt	acaagatcat	tgttatctga	regeeeeeee	agacttttta
27901 27961	cotttaccac	aatgctgata cccgtttcca	aattgacctc	catactata	taatttatct	gaccacacac
28021	gatttaaac	caaatcaggt	cactttaacc	acactattee	totcaatctt	taccaattca
28081	ctattgagcc	tatttccctc	gcccacctc	tattggttgc	tacctatttt	tctttcatt
28141	cacataactc	tgaatgccat	tgatggcatg	ctggcacggg	aacataacca	gaagtctcat
28201	ctgggcgcta	tttataatga	attgggggat	gtcatttctg	atgttgccct	ctacctcccc
28261	ttctgccttt	tacctgatgt	gaacagcctc	agcctgttga	ttattttatt	cctcactatc
28321	ttgaccgaat	tcatcggcgt	actggcacaa	acgattggtg	catcacggcg	ctatgacggc
28381	ccgataggaa	aaagtgaccg	tgcttttatc	ttcggagctt	atggattgat	tattgcgatt
28441	ttccctttgg	ccttgggctg	gagtatetet	ttgtttgctt	tcatgatcat	tttactcttg
28501	gtgacttgct	atcagcgcgt	tgttaaagcc	ttacgtgaaa	teeggetgge	tgaacagtca
28561	cactccaaat	gaggcgttaa catcagataa	catgacacca	ttttaggett	actuactige	acangaacac
28621 28681						acgtatccag
28741	catataatta	acquactcat	tetacetaat	attectatet	tcacataaga	tgcccgtgga
28801	cacgotgaga	cagaagggc	gcgcggttac	agcccatcca	tgggaacgtc	gattcgtgat
28861	ottoatoaat	ttotcagatt	tattgccact	cagtacggca	tcgccatgga	aaatatcgtg
28921	gttatcggcc	agagtgtcgg	agcggtatta	gtctctgctt	gggtacacga	ctatgcgcca
28981	aaaatccgcg	ccatgatcct	cgcagcaccc	gcatttgata	ttaaattgta	tatccctttt
29041	gccacgcagg	gactgcaatt	gatgcaaaaa	gcacgaggta	ttttcttcgt	gaattcctat
29101	gtgaaagcca	gatatctgac	tcacgatgaa	acccgaattg	cctcttataa	tagcgatccg
29161	ttgattaccc	gggaaatcgc	cgtcaatatt	ctcttggatc	tttaccaaac	cgccgagcga
29221	gtagttaaag	atgccgccgc	cattacacta	cctaccctgt	tgtttattte	aggcagcgat
29281	tatgtagtga	acaaaaaacc	acagcatcag	ttttatcagc	agctaaatac	ccctatcaaa
29341	gaaaaacatg	tgatggatgg	CTTCTACCAC	gatacgttgg	gradadaga	tcgccatctg
29401	gtttttgaca	aaatccgggt	ctcattgag	tetaceate	aatttcaasa	ttatcagcac attaagcaca
29461 29521	trattarret	atctatata	taagaaacto	anctatoast	taatacataa	ggtaatgagt
29521 29581	actoactor	graduactto	caagaaaccc	tocatcootc	tcaaaarnnn	gtttgattcc
29561						tttggggcga
29701	atactcoata	agcattattt	gaacagcatt	gattaacaca	gtatacgcca	gcgcaagatc
29761						tatgcctgtg
29821						caacgatttc
				2/1/1	-	,

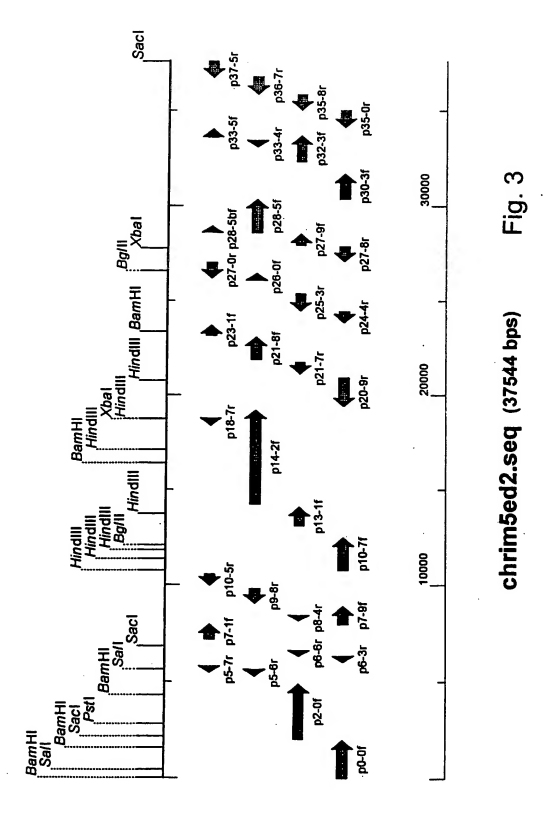
Fig. 2(vi)

chrim5	ed2.seq					
29881	agcaaagtcg	attctatttt	gttaagggac	tatagcgaaa	tcaatgttaa	tcaagggcag
29941	gcttatattg	aggagcgcga	tctgacggac	aaaattcgtt	ttattatcgg	tgatgccttt
30001	aatgctgaaa	gcatctcatc	cattacgcca	gcgccgacac	tgggtattgt	atccggtctc
30061	tatgaattgt	tccctgataa	taatttactc	agaaattcgc	tacgcggctt	tgctgatgtt
30121	atgacagaaa	atggttatct	ggtgtacacc	ggccaaccgt	ggcatccaca	aactgaggtc
30181 30241	ategeeegtg	tageageett	ecategigae	agecaacege	ggatcatgcg aaaaactgta	ccaactgaca
30301	cataactaga	ggattttcac	tatttcaatt	accaaacata	ttcatcgctg	atgaataaat
30361	aaatataaga	tggaacacca	catgcactct	tctctcgata	gtcgtcggcg	cctatggctg
30421	acaggtgtta	tctggctatt	gtttctggct	ccgtttttct	ttcttactta	tggccaggtc
30481	aatcagttca	cggcacaaag	aagcgatgtc	ggcactgtga	tgttcggttg	ggaacataac
30541	atccctttt	ggtcatggtc	gattatccct	tactggagta	tcgatctgtt	ctacggaata
30601	tcgttattta	tctgtaccca	tcgccgtgaa	cagtggcttc	acggctggcg tgaaattttc	attattaca
30661 30721	ccaccacca	aaggeetatt	taactaatta	tttaatcaac	tggagttatt	tgatctgcc
30781	tataatcaag	ccccttccct	gcacattatt	ctactataat	tgctctggct	gcgctattca
30841	gcctacgtga	gtggttactg	gcgtgggttg	ctgcacattt	ggtcagtgct	gattgcactc
30901	tcggttctga	cgacttggca	gcaccatttt	atcgatgtac	taacgggttt	tgccgttggt
30961	gtcatcctca	gttacctact	gccggtttca	taccgctggc	gctggcaacc	taatcaagat
31021	cgctatgcac	ggaagttatt	cggctattat	ctgacaggca	gcgctttgtt	cgcgcttata
31081 31141	gcgagtctgc	cctacacac	attaggeage	tecatattte	ctgctgtatc aaaaacagcc	agatogcog
31201	atotcactot	ctgcacgctg	gctactggcg	ccataccaac	tgggagcatg	gctctcttat
31261	ctctggttcc	ggcgtaaaag	cgcacctttc	aaccatataa	ctgaagggat	tattctcggc
31321	agcctgcctt	gccagcccgt	tacggcggtc	agtgtccttg.	atataaccgc	tgagtggcac
31381	aggcgatcgg	atgcccgcac	agtaaattat	gtttgccagc	cgcaaatcga	cttactgccg
31441 31501					ataaactacg cgatggtggt	
31561	ctactgaaac	agcatcctga	atatgatate	aacactgtcg	tagcaatcct	gcgtaaagcc
31621	agaccgcatg	tcacgttcag	acaaacacat	ctggatgccc	tgtctcaatg	ggcaaaaggc
31681	tacctataac	ggggaacata	acatgcagcc	ggaaaatctg	atcagcaaag	tgattatcgc
31741					ttttctatcc	
31801	tgcaatgctg	attgctgtct	tcaacactac	cgccttaaac	aacattgcac cgcacttggc	ttgactgccgt
31861 31921	ctattttcad	atcctcaatt	catecectea	aaaaagggg	gagttcgatc	aaacottatt
31981	gctgatattt	aacaagttac	cccaatcacg	gacacagaat	gatcgcttta	acggagcaat
32041	caaactgtta	aaaaaggcta	cgattggtct	gatcctgcaa	tggatactgt	ttttcctgtt
32101	tetgettact	ttgaaatatt	cagcctgaat	tcagctttaa	tctgacaaag	tcagaacgta
32161	acgatgctac	attttctcga	ctgtactact	attaactaac	ttttagacct	graacacett
32221 32281	tattgggtga	ttttagatta	actettacat	cataasatta	agcccgtcat atgggatacg	cccatttagt
32341	gcgtttattg	actcatqtct	gaaggaatct	tactctttcc	cccgttttat	cagagatatc
32401	atcgctggga	ttaccgtggg	tgttatcgct	atcccattgg	caatggctct	ggcgatagga
32461	agcggagttg	caccacagta	tggcctgtat	actgccgcta	tegeeggeat	tgtcatcgcc
32521	atgaccggag	gctcacgtta	tagcgtttcc	ggcccaacgg	cggcttttgt	ggtgatcctt
32581 32641	ratectgttt	ttatastaaa	attaccacat	tttaascaac	ttgcgacgct. ttatcgaata	tattcctato
32701	tetettacce	ttggatttac	ttccggtatt	accatcacta	tcgctaccat	gcaggtgcaa
32761	aatttctttg	gcctgaaact	ggcacatata	ccggaaaatt	atattgataa	ggtagttgca
32821	ctttatcaag	ccctgccttc	attacaattg	agtgatacgc	ttatcgggct	gactacgctt
32881	ctggtactga	ttttctggcc	gaaactggga	gtaaagttac	cgggtcactt	accogctttg
32941	atcgcgggta	cagctgttat	gggcgcaatg	catctgctga	accatgatgt	tocaccatt
33001 33061	cttccccat	ttatcctacc	accygeggat	cccatacac	gtcaaggcat attctctcga	tattagctgg
33121	aataccgtat	cggcattgct	acccactaca	ttttcgatgg	ccatgctggg	tgcgattgaa
33181	tcgttactgt	gtgcagtgat	tctggatggt	atgacaggga	aaaaacacca	ttctaacggt
33241	gagetgettg	gccaggggtt	gggcaatatt	gccgcccctt	tetttggtgg	cattaccgcc
33301	actgcggcta	tegecegite	agccgccaat	gtgcgggcag	gtgcaacttc	cccgatagcc
33361 33421	getgtggtge	tagagagant	ggtattatta	acattgetgg	ttttggctcc tcgcgtggaa	tatgaccaa
33481	descataadd	tagtggattt	aataccccat	gcgcctaaag	atgacattat	tatcatactt
33541	ttatatctat	cattgactgt	cctattcgat	atggtccgtc	gcgatcacta	tcggcattgt
33601	gctggcatca	ctcctgttta	tgcgcaaaat	tgccaatatg	actcgaatca	gcacgtcatc
33661	tttaacaagc	gcggagaaag	ggttattggt	cgtacgaatt	aacggccctt	tattcttcgc
33721	tgccgccgaa	cgtatttttg	ccgaactgag	agaaaaaagt	gctgattatc	aaaccatcat
33781	catgcagtgg	gatgccgtac	ccgttttgga	cgctggcgga	ttacacgctt	ttcagggttt
33841 33901	tgtgcgcgaa	cccggcaaag	aaaaacatat	cgtcgtatgt	gatattccct	tccagccatt tttatgctac
33961	cttacccaa	gcactasaa	agatagee	tgattacacg	cctgaggtc	gtgcttcttc
34021	agaaaagatt	caaggtcagt	aacaacaagt	caccctctoc	cagttatoct	ggcagagggt
34081						gaaattttc
34141						caatggcaaa
			Eia 2/	۸ii۱		

chrim5ed2.seq gcagacgggc catcacattt ggcattttct ggatctgggt gcgcttcaag gaacagacct gcgatccca cagccatacc cgcacgagcc agttcagcaa cctgcgcgcg acggccaccg 34261 34321 gaagetgcac ctaatgggte geggeactgt aatgeatgag tgaegtegaa aatgaeegge gcaccttggg tagcttgttg catgacaccg aagcccagca tgtcaaccac cagattgtca taaccaaagt tgctgccacg gtcacacagg atcacctggt cattgccgc ttctttgaat ttttcaacaa tattgcccat ttgcccaggg ctaacaaaact gtggttttt aacattaatc 34381 34441 34501 34561 acagcaccgg tittegecat egetteaace agateggtet ggegggeaag gaaggetgga 34621 agetgaatea catecaceae atecgegaet ggttgageet gtgcaggete atgcaegtea gtgataattt teaegeegaa ggtttgttte agtteetgaa agatttteat eeeetetee 34681 agecetggee cacgataaga geggatagaa gaaeggttag cettateaaa agaegettta aaaacataag gaatgeecag tttttgtgte aeggtgaeat agtgeteaca aateegeatg 34741 34801 34861 gccaaatccc gtgattcaag gacgttcatt ccaccaaaca atacaaatgg cagatcattt 34921 gegaeettga tateaceaat attaaceaet ttetgttgea tgettataee ttetettgtt aagggcaaat aaatttgctg tatcaataaa aaaataaacc taatgcagga caatcaaccg 34981 35041 ctgttctatg gtgttaattt gcatttttat catttctgag attgggtcct cagggcaatg 35101 ttcgacaaaa taactcaaat ccgaaacagc aatatgattg cagtcgagct gggcatagat aageeeeega tegeggattt catagggate ategggatea aaegteagea eeaetteget 35161 35221 ggeettaage geeagtteea tettitete tteeateaga gaaaettiga tggtgteagt 35281 gattttgcgc acgaggctga cattatcggc ttcctgcaaa tcctgttttt tcagacgaac 35341 agttgggcca atattacctt ttaaccagac atccagcgta tgttcattca gcgtatcccc 35401 attcaaggga ttaatgaacc aagtgggctg atcaagcaaa tcaatccgca atatcagttg 35461 gattggaaat atgacagget gtacegacag ecceagegee tgagcaatat gegtaaacae cgtacccaat gatacaggtg agccttgacg tgaatggagt aagcgatcca gccacagggt atctgataga caatacaccc cattagctcc accaaacttc cattcccgat aaaaaagtgt 35521 35581 tagcagcgaa tccaatttca ctttggaatc ggtaatggag gaaagcctct gccgggcttc ttcaaccaat gcagttagct gctcactcac cagagtctga ggaaaatcag ggcggataat ttttgatacc agaataatac cgtcaatcag gggagtatta ttgaattcat aattaactat 35641 35701 35761 35821 gggtitcatt titattccat cgacctatcg tgatgcgttc attaccgcca taatcctgaa aagtcgctat ctgttgataa cccttctcta aaaataggtt tctgacaacg gttccctgtt 35881 tccagccatg ttccagcaat agccatccat ttggtgacag gaaatggcgc gcctgtccca caattgcctg caaatccgcc atgccattt gtgcagcgat caatgcagtg gctggttcaa acctgatatc cccttcttgt agatgaggat cacgctcatc tatatacgga ggattgctga 35941 36001 36061 caatcatatc aaattgttgg ttacccactg ctgcaaacca ctcactttgc aaaaaattca cattgtgaat ggccagtttt ccggcgtttt tttcagcatt gtgttgtgcc agcatcacgg catcagagtt gatatcgacc cctgtcacat aacaatcatt ccgctcactt gccaatgcca 36121 36181 36241 36301 gtgcaatcgc ccccgtcccc gtccccagat ccagaatccg ggctggagaa tcaggcaata 36361 attccaatgc cttctccacc agacattcag tatcagggcg cgggatcaac gtcgctggcg atacggcaaa cggcagtgac cagaattccc gttcaccaat aatataagct accggctctc 36421 cctgaatgcg gcgcaccagc aggctatcaa gctgatgcaa ttcttccgat gagattagcg tttcatcgaa agcaatcaga taagtacggg aacgccctgt cacgtatccc aacaagattt ccgcatcacg cttagggctg tcactttcag acaactgggt agccgcatgt tgtagccagc 36481 36541 36601 36661 attggtaatt cattaatcct gctctgacag tgcagacagt tgatctgcct gatattcgat aatgatcggc tgaataagca tatccagttt gccttctatc acttcatcaa ggcggtataa 36721 36781 cgtcagattg attcggtgat cagtcacacg cccctgtggg aagttatagg tgcggttgcg atctgagegg teacagaac cagcaactt teggegttea gaggettea ettettggeg etteegeate teageageac ggatacaget tgecaatacg gacategett ttgetttgtt ttgtgetgg gaacgeteat cetgacatte cactacgate eccepttgga gatgggtaat tegaatege gaateggtg tattgaegtg etgecaacac geaceggaag agegaaacgt atctattte aaateaceeg ggetgatgte eggeatetea getteetggaa ttetetggaa 36841 36901 36961 37021 37081 37141 gacagccaca gtacaggcag aagtgtgaat gcgcccctga gattccgttt ccggtacacg 37201 ctggacacga tgaccgcctg attcaaattt caagtgacca taaacctgat cacccgaaac triggeaate actientity agreeaceaty etergeeticy triggegetta taateetaate tetecagegg catagegget atacatgegg aacaaatete eggatette tegecacegg treetgeegg gaetteaagg aacaaatete eggatette ggeaacagea gtagetgtag etgetgtee agetetteat taegaattit tgeeteettg agetetteet gegeeatte eeggatette 37261 37321 37381 37441 37501

Fig. 2(viii)

11/14



SUBSTITUTE SHEET (RULE 26)

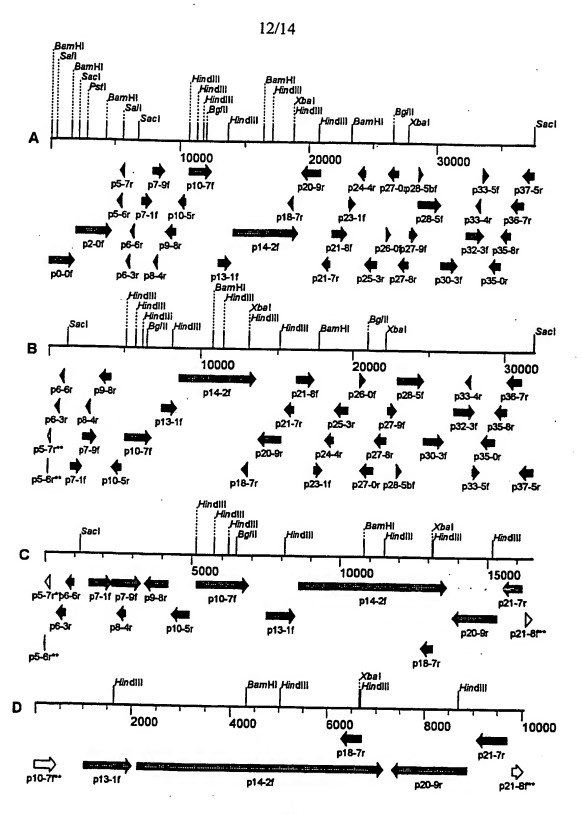
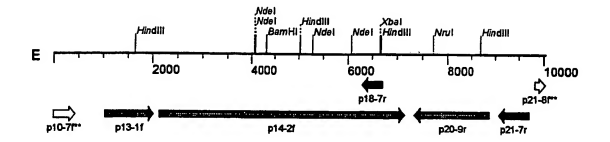
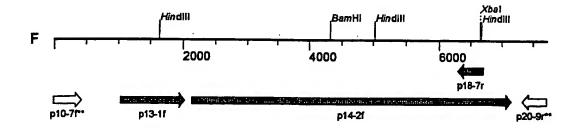
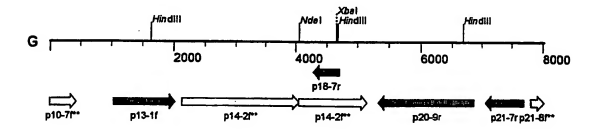


Fig. 4







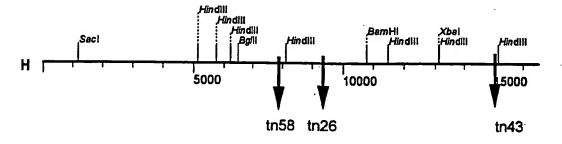
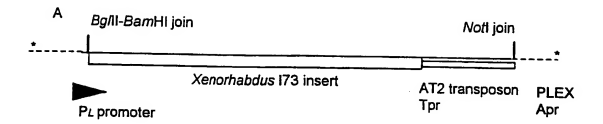


Fig. 4(cont'd)

14/14



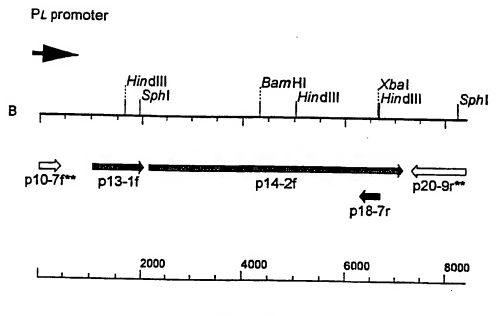


Fig. 5

int. onal Application No PCT/GB 00/00219

		101/40 00/00219
A CLASSIF IPC 7	CATION OF SUBJECT MATTER A01N63/00 A01N63/02 C12N15, //(C12P21/00,C12R1:01)	/31 C12P21/00 C07K14/24
According to	International Patent Classification (IPC) or to both national classification	fication and IPC
B. FIELDS		
Minimum doc IPC 7	oumentation searched (classification system followed by classific AOIN	ation symbols)
Documentati	on searched other than minimum documentation to the extent the	t such documents are included in the fields searched
Electronic de	ata base consulted during the international search (name of data	base and, where practical, search terms used)
C. DOCUME	ENTS CONSIDERED TO BE RELEVANT	
Category *	Citation of document, with indication, where appropriate, of the	relevant passages Relevant to claim No.
P,X	WO 99 22598 A (UNIV READING ;EL ABDULRAHAMAN (GB); HAGUE NIGEL 14 May 1999 (1999-05-14) cited in the application page 1 -page 2 page 4, line 7 - line 14	
	page 5, line 2 - line 8 page 5, line 25 - line 28 page 8, line 1 - line 15; claim 1-3,9,10,14; example 5	s -/
	ner documents are listed in the continuation of box C.	Y Patent family members are listed in annex.
X Furth	ner documents are listed in the continuation of box C.	X Paton farmy fromboto are union.
"A" docume consid "E" earlier of filing d "L" docume which citation other "P" docume other "P" docume	tegories of cited documents:  ant defining the general state of the art which is not leved to be of particular relevance socument but published on or after the international late and which may throw doubts on priority claim(a) or is cited to establish the publication date of another nor other special reason (as specified) ent referring to an oral disclosure, use, exhibition or means ent published prior to the international filing date but han the priority date claimed	"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person eldiled in the art.  "&" document member of the same patent family
	actual completion of the international search	Date of mailing of the international search report
1	6 May 2000	13/06/2000
Name and r	mailing address of the ISA  European Patent Office, P.B. 5818 Patentlaan 2 NI. – 2280 HV Rijawijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	Authorized officer  Muellners, W

1

triti anal Application No PCT/GB 00/00219

C/Continu	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	PCT/GB 00/00219
Category *		Industrial Alexander
	the second and a second	Relevant to claim No.
X,P	DATABASE BIOSIS 'Online! BIOSCIENCES INFORMATION SERVICE, PHILADELPHIA, PA, US AN 2000:109497, GREWAL, PARWINDER S. (1) ET AL: "Allelopathy: A possible mechanism of suppression of plant-parasitic nematodes by entomopathogenic nematodes." retrieved from STN XP002136524 abstract & NEMATOLOGY, (NOV., 1999) VOL. 1, NO. 7-8, PP. 735-743.	1-9,11,
X,P	DATABASE BIOSIS 'Online! BIOSCIENCES INFORMATION SERVICE, PHILADELPHIA, PA, US AN 2000:109391, HAN, RICHOU ET AL: "Trans - specific nematicidal activity of Photorhabdus luminescens." retrieved from STN XP002136525 abstract & NEMATOLOGY, (NOV., 1999) VOL. 1, NO. 7-8, PP. 687-693.	1-9,11, 29
X,P	CHEMICAL ABSTRACTS, vol. 132, Columbus, Ohio, US; abstract no. 163304, HU, KAIJI ET AL: "Nematicidal metabolites produced by Photorhabdus luminescens (Enterobacteriaceae), bacterial symbiont of entomopathogenic nematodes" XP002136522 abstract & NEMATOLOGY (1999), 1(5), 457-469,	1-9,11, 29
X,P	WO 99 42589 A (NOVARTIS ERFINDUNGEN VERWALTUN; NOVARTIS AG (CH); KRAMER VANCE CAR) 26 August 1999 (1999-08-26) cited in the application page 1 -page 2, paragraph 3; claims	12-18
X	CHEMICAL ABSTRACTS, vol. 126, Columbus, Ohio, US; abstract no. 234686, GEORGIS, R. ET AL: "Novel pesticidal substances from the entomopathogenic nematode-bacterium complex" XP002136523 abstract & ACS SYMP. SER. (1997), 658(PHYTOCHEMICALS FOR PEST CONTROL), 134-143,	1-9,11,

Int. onal Application No PCT/GB 00/00219

	PCT/GB 00/	0/00219				
	(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT					
Category *	Citation of document, with indication, where appropriate, of the relevant passages		Relevant to daim No.			
X	K. HU ET AL.: "Mortality of Plant-Parastic Nematodes Caused by Bacterial (Xenorhabdus spp. and Photorhabdus Luminescens) Culture Media" JOURNAL OF NEMATOLOGY, vol. 27, no. 4, December 1995 (1995-12), XP000905673 the whole document		1-9,11, 29			
X	WO 98 08388 A (MORGAN JAMES ALUN WYNNE; JARRETT PAUL (GB); ELLIS DEBORAH JUNE (GB) 5 March 1998 (1998-03-05) cited in the application page 1, line 4 - line 9 page 6, line 21 -page 9, line 9 page 11, line 10 -page 12, line 19; claims 17,18,21,23-25,29	·	6-9, 11-30			
X	PATENT ABSTRACTS OF JAPAN vol. 013, no. 286 (C-613), 29 June 1989 (1989-06-29) -& JP 01 080294 A (SUMITOMO CHEM CO LTD;0THERS: 01), 27 March 1989 (1989-03-27) abstract; figures 2-1		12–18			
A	WO 92 19739 A (MYCOGEN CORP) 12 November 1992 (1992-11-12) cited in the application		1-30			
X	page 1 -page 2, line 32; claims		12-18			

....mational application No.

PCT/GB 00/00219

Box I	Observations where certain claims were found unsearchable (Continuation of Item 1 of first sheet)
This Inte	emational Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1.	Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:
2. 🗌	Claims Nos.:  because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically:
з. 🗌	Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II	Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This Inter	mational Searching Authority found multiple inventions in this international application, as follows:
1. 🔲	As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.
2	As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3	As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:
4 n	No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is estricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark o	The additional search fees were accompanied by the applicant's protest.  No protest accompanied the payment of additional search fees.

information on patent family members

Intic onal Application No PCT/GB 00/00219

	atent document d in search report		Publication date		Patent family member(s)	Publication date
WO	9922598	A	14-05-1999	AU	9753098 A	24-05-1999
WO	9942589	A	26-08-1999	AU	3028699 A	06-09-1999
WO	9808388	A	05-03-1998	AU	4024997 A	19-03-1998
•••				CN	1233938 A	03-11-1999
				EP	0923295 A	23-06-1999
				ZA	9707373 A	15-02-1999
JP	01080294	Α	27-03-1989	JP	2117839 C	06-12-1996
٠.	01000	••		JP	8017707 B	28-02-1996
WO	9219739	A	12-11-1992	AU	667041 B	07-03-1996
**-		• •		AU	2025292 A	21-12-1992
				AU	656726 B	16-02-1995
				AU	7826491 A	12-12-1991
				BR	9205969 A	26-07-1994
				CA	2042868 A	12-12-1991
				EP	0462721 A	27-12-1991
				EP	0517367 A	09-12-1992
				HU	62928 A	28-06-1993
		•		JP	4229170 A	18-08-1992
				NZ	242560 A	26-01-1994
				PL	290626 A	24-02-1992
				ÜS	5439881 A	08-08-1995
				ÜS	5753492 A	19-05-1998
				ZA	9203168 A	27-01-1993

# THIS PAGE BLANK (USPTO)